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FORM PTO-1390 (REV. 11-2000)	US DEPARTMENT OF COM	IMERCE PATENT AND TRADEMARK OFFICE	ATTORNEY 'S DOCKET NUMBER
	ANSMITTAL LETTER	TO THE UNITED STATES	10365/07304
		ED OFFICE (DO/EO/US)	U.S. APPLICATION NO (If known, see 37 CFR 15
		NG UNDER 35 U.S.C. 371	Unassi 199/937052
INTERNA	TIONAL APPLICATION NO.	INTERNATIONAL FILING DATE	PRIORITY DATE CLAIMED
PCT/US	00/07995	23 March 2000 (23.03.00)	23 March 1999 (23.03.99)
- f-	INVENTION		
	BETA-TUBULIN GENES NT(S) FOR DO/EO/US		
	er S. SIDHU, et al.		
Applicant l	nerewith submits to the United Sta	ates Designated/Elected Office (DO/EO/US)	the following items and other information
1. XX Th	s is a FIRST submission of items	s concerning a filing under 35 U.S.C. 371.	
2. Th	s is a SECOND or SUBSEQUE	NT submission of items concerning a filing t	ınder 35 U.S.C. 371.
ite	ns (5), (6), (9) and (21) indicated		
		ration of 19 months from the priority date (A ion as filed (35 U.S.C. 371(c)(2))	article 31).
5. XX A c	<u> </u>	only if not communicated by the Internation	nal Rureau)
b.	has been communicated by	·	Mai Duleddy.
c,		ication was filed in the United States Receiv	ing Office (RO/US).
6. An	English language translation of the	ne International Application as filed (35 U.S.	.C. 371(c)(2)).
a.	is attached hereto.		
b.		tted under 35 U.S.C. 154(d)(4).	
7. XX Am		ernational Aplication under PCT Article 19 (* * * * * *
a.		ed only if not communicated by the Internati	onal Bureau).
b.	have been communicated b	by the International Bureau.	
c.	have not been made; howe	ver, the time limit for making such amendme	ents has NOT expired.
d.	have not been made and w	ill not be made.	
8. An	English language translation of the	e amendments to the claims under PCT Arti	cle 19 (35 U.S.C. 371 (c)(3)).
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	English lanugage translation of thicle 36 (35 U.S.C. 371(c)(5)).	ne annexes of the International Preliminary E	Examination Report under PCT
Items 1	to 20 below concern document	t(s) or information included:	
11. 🔲 A	n Information Disclosure Stateme	ent under 37 CFR 1.97 and 1.98.	
12. 🔲 A	n assignment document for record	ding. A separate cover sheet in compliance	with 37 CFR 3.28 and 3.31 is included.
13. A	FIRST preliminary amendment.		•
14. 🔲 A	SECOND or SUBSEQUENT pr	eliminary amendment.	
15. A	substitute specification.		
16. 🔀 A	change of power of attorney and	or address letter.	
17. XX A	computer-readable form of the se	equence listing in accordance with PCT Rule	13ter.2 and 35 U.S.C. 1.821 - 1.825.
18. A	second copy of the published into	ernational application under 35 U.S.C. 154(d	1)(4).
19. 🗌 A	second copy of the English langu	age translation of the international application	on under 35 U.S.C. 154(d)(4).
20. XX O	her items or information: Sta	tement under WIPO Standard S	r.25
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4	e and is addressed to the Assistant Com-		

Derrick Gordon (Name of Depositor)

Date of Signature: 19 September 2001

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21. XX The following fees are submitted:					CAL	CULATIONS I	TO USE ONLY
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Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO							
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d. Fees are to be charged to a credit card. WARNING: Information on this form may become public. Credit card							
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NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137 (a) or (b)) must be filed and granted to restore the application to pending status.							
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FUNGAL BETA-TUBULIN GENES

TECHNICAL FIELD OF THE INVENTION

The invention relates to a bio-affecting composition and to a biological diagnostic and experimental agent.

BACKGROUND OF THE INVENTION

TAXOL® (Bristol-Myers-Squibb), generically known as paclitaxel (hereinafter referred to as "taxol"), is a complex diterpenoid which has demonstrated anti-tumor activity against breast and ovarian cancer (Rowinsky, E.K. and Donehower, R.C. 1991. "The clinical pharmacology and use of antimicrotubule agents in cancer chemotherapeutics," Pharmacol Ther 52:35-84). Its anti-tumor activity is due to its ability to bind to beta-tubulin in assembled microtubules (MTs) and stabilize them (Manfredi, J.J. and Horwitz, S.B. 1984. "Taxol: an antimitotic agent with a new mechanism of action," Pharmacol Ther 25:83-125; and Horwitz, S.B. 1992. "Mechanism of action of taxol," Trends Pharmacol Sci 13:134-136). In vivo, taxol affects spindle function during mitosis, resulting in cell cycle arrest in G2/M phase. In vitro, taxol promotes MT assembly and prevents their disassembly under conditions which would otherwise cause depolymerization (Schiff, et al. 1979. "Promotion of microtubule assembly in vitro by taxol" Nature 277:665-667; and Pamess, J. and Horwitz, S.B. 1981 "Taxol binds to polymerized tubulin in vitro," J Cell Biol 91:479-487). The taxol binding site on beta-tubulin has been characterized by photo cross-labeling, electron crystallography, and mutagenesis, and involves several regions of beta-tubulin (Rao, et al. 1994. "3'-(p-Azidobenzamido) taxol photolabels the N-terminal 31 amino acids of βtubulin," J Biol Chem 269:3132-3134; Rao, et al. 1995. "Characterization of the taxol binding site on the microtubule," J Biol Chem 270:20235-20238; Nogales, et al. 1998. "Structure of the aß tubulin dimer by electron crystallography," Nature 391:199-203; Nogales, et al. 1999. "High-resolution model of the microtubule," Cell 95:79-88; and Giannakakou, et al. 1997. "Paclitaxel-resistant human ovarian cancer cells have mutant βtubulins that exhibit impaired paclitaxel-driven polymerization," J Biol Chem 272:17118-17125).

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Taxol was found originally in the inner bark of pacific yew trees (Taxus brevifolia) by Wani et al. (Wani, et al. 1971. "Plant antitumor agents. VI. The isolation and structure of taxol, a novel antileukemic and antitumor agent from Taxus brevifolia," J Am Chem Soc 93:2325-2327), and noted to constitute about 0.02% of dry phloem weight. The limited resource of yew trees made it advantageous to locate additional sources for taxol.

In 1993, Stierle et al. reported the isolation of a taxol-producing fungus, Taxomyces andreanae, an endophyte associated with T. brevifolia (Stierle, et al. 1993. "Taxol and taxane production by Taxomyces andreanae, an endophytic fungus of pacific yew," Science 260:214-216). T. andreanae produces authentic taxol, though at very low levels (24-50 ng/liter of liquid culture). Recently, additional taxol-producing fungi have been isolated, including various strains of Pestalotiopsis microspora (Li, et al. 1996. "Endophytic taxolproducing fungi from bald cypress, Taxodium distichum, "Microbiolog 142:223-226; Li, et al. 1998. "The induction of taxol production in the endophytic fungus-Periconia sp. from Torreya grandifolia," J Ind Microbiol Biotechnol 20:259-264; Strobel, et al. 1996. "Taxol from fungal endophytes and the issue of biodiversity," J Ind Microbiol 17:417-423; and Strobel, et al. 1996. "Taxol from Pestalotiopsis microspora, an endophytic fungus of Taxus wallachiana," Microbiolog 142:435-440). One P. microspora strain, Ne32, was isolated from the inner bark of Himalayan yew T. wallachiana, and produces approximately 50 µg taxol per liter of liquid culture (Strobel, et al. 1996. Microbiolog 142:435-440; and Li, et al. 1998. "Stimulation of taxol production in liquid culture of Pestalotiopsis microspora," Mycol Res 102:461-464). The taxol produced from fungal sources has been reported to be spectroscopically and chromatographically identical to taxol isolated from yew trees, and has shown similar pharmacological effects on cancer cell lines (Strobel, et al. 1996. Microbiolog 142:435-440). The production of taxol from fungal sources has provided broader resources of taxol, reduced production costs, and a means of meeting the increasing demand for taxol.

While taxol has been shown to be toxic to cells from a wide range of organisms including mammals, sea urchin, *Drosophila, Xenopus, Physarum, Haemanthus*, and *Trypanosoma* (Baum, et al. 1981. "Taxol, a microtubule stabilizing agent, blocks the replication of *Trypanosoma cruzi*," *Proc Natl Acad Sci USA* 78:4571-4575; Kellogg, et al. 1989. "Proteins in the centrosome, spindle, and kinetochore of the early *Drosophila*

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embryo, "J Cell Biol 109:2977-2991; and Manfredi, J.J. and Horwitz, S.B. 1984. Pharmacol Ther 25:83-125), variable sensitivity to taxol has been reported in fungi. Young et al. tested taxol toxicity on representative species from different fungal groups (Young, et al. 1992. "Antifungal properties of taxol and various analogues," Experientia 48:882-885). In Young's study, five oomycete species were identified as sensitive to taxol (IC₅₀ 0.4-5.9 μ M), including the plant pathogens Pythium ultimum and Phytophthora capsici. In P. capsici, taxol inhibited nuclear division at low concentrations, indicating that it acts through a mechanism similar to that in mammalian cells. In contrast, four ascomycete species were identified as resistant to taxol (IC₅₀ > 50 μ M). This resistance was reported to be due to the reduced ability of fungal microtubules to interact with taxol. Taxol was also shown to be unable to stabilize MTs assembled with purified S. cerevisiae tubulin (Bames, et al. 1992. "Yeast proteins associated with microtubules in vitro and in vivo," Mol Biol Cell 3:29-47) and only weakly stabilize MTs from Aspergillus nichulans (Yoon, Y. and Oakley, B.R. 1995. "Purification and characterization of assembly-competent tubulin from Aspergillus nichulans," Biochem 34:6373-6381).

Because of the anticancer properties of taxol and the variability of fungi to taxol sensitivity, there is a continuing need for isolating and/or identifying novel beta-tubulin genes useful for developing isogenic fungal strains that are either taxol-sensitive or taxol-resistant. These beta-tubulin genes and/or isogenic fungal strains can then be applied to anticancer drug screening and for developing diagnostic tests for tumor sensitivity assays.

SUMMARY OF THE INVENTION

In one aspect, the invention is a purified DNA segment encoding a beta-tubulin of the fungal species *Pestalotiopsis microspora* or a portion thereof. Preferably, the DNA segment encodes at least one taxol binding site. For some uses, it is preferable that the DNA segment encodes a protein having taxol binding site I and taxol binding site II. For DNA segments that encode proteins which function as beta-tubulins, the DNA segment encodes a protein which has taxol binding site I and taxol binding site II and is able to interact with alpha-tubulin. An exemplary DNA segment comprises at least a portion of SEQ ID NO:1. Another exemplary DNA segment comprises a portion of SEQ ID NO:1, with or without substitution. Another exemplary DNA segment comprises a portion of SEQ ID NO:1, with or

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NO:1 comprising the nucleotide sequence from nucleotide 708 through nucleotide 764 of SEQ ID NO:1, with or without substitution. Another exemplary DNA segment comprises a portion of SEQ ID NO:1 comprising the nucleotide sequence from nucleotide 708 through nucleotide 764 of SEQ ID NO:1, wherein either nucleotide 729, nucleotide 730 or nucleotide 731 or mixtures thereof are substituted. Another exemplary DNA segment comprises the nucleotide sequence from nucleotide 75 to nucleotide 1412 of SEQ ID NO:1 wherein the DNA segment encodes a beta-tubulin. Another exemplary DNA segment comprises the nucleotide sequence from nucleotide 75 to nucleotide 1412 of SEQ ID NO:1 wherein at least one nucleotide in the nucleotide sequence is substituted and wherein the taxol binding capacity of the beta-tubulin is not altered. Another exemplary DNA segment comprises the nucleotide sequence from nucleotide 75 to nucleotide 1412 of SEQ ID NO:1 wherein at least one nucleotide sequence from nucleotide 75 to nucleotide 1412 of SEQ ID NO:1 wherein at least one nucleotide in the nucleotide sequence is substituted and wherein the taxol binding capacity of the beta-tubulin is altered.

In another aspect, the invention is an amino acid sequence comprising at least a portion of a beta-tubulin of the fungal species Pestalotiopsis microspora. Preferably, the amino acid sequence comprises at least one taxol binding site. For some uses, it is preferable that the amino acid sequence is a protein having taxol binding site I and taxol binding site II. For amino acid sequences that can function as beta-tubulins, the amino acid sequence has taxol binding site I and taxol binding site II and is able to interact with alphatubulin. An exemplary amino acid sequence comprises at least a portion of the beta-tubulin as depicted in SEQ ID NO:2. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:2 comprising Amino Acids 1-31, with or without substitution. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:2 comprising Amino Acids 212-230, with or without substitution. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:2 comprising Amino Acids 212-230 with an amino acid substitution at Amino Acid 219. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:2 consisting essentially of Amino Acids 1-446 wherein the portion behaves as a taxol-resistant beta-tubulin. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:2 consisting essentially of Amino Acids 1-446 the portion contains at least one amino acid substitution that alters the taxol binding property of the portion. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:2 consisting essentially of Amino Acids 1-446 the portion contains at least one amino acid

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substitution that does not alters the taxol binding property of the portion. Another exemplary amino acid sequence is substituted with any amino acid which perturbs the three-dimensional structure of the amino acid sequence surrounding Amino Acid 219 as numbered in SEQ ID NO:2.

In another aspect, the invention is a purified DNA segment encoding a beta-tubulin of the fungal species Pythium ultimum or a portion thereof. Preferably, the DNA segment encodes at least one taxol binding site. For some uses, it is preferable that the DNA segment encodes a protein having taxol binding site I and taxol binding site II. For DNA segments that encode proteins which function as beta-tubulins, the DNA segment encodes a protein which has taxol binding site I and taxol binding site II and is able to interact with alphatubulin. An exemplary DNA segment comprises at least a portion of SEQ ID NO:3. Another exemplary DNA segment comprises a portion of SEQ ID NO:3 comprising nucleotide 92 through nucleotide 184, with or without substitution. Another exemplary DNA segment comprises a portion of SEQ ID NO:3 comprising the nucleotide sequence from nucleotide 725 through nucleotide 781, with or without substitution. Another exemplary DNA segment comprises a portion of SEQ ID NO:3 comprising the nucleotide sequence from nucleotide 725 through nucleotide 781, wherein either nucleotide 746, nucleotide 747 or nucleotide 748 or mixtures thereof are substituted. Another exemplary DNA segment comprises the nucleotide sequence from nucleotide 92 to nucleotide 1429 of SEQ ID NO:3, wherein the DNA segment encodes a beta-tubulin. Another exemplary DNA segment comprises the nucleotide sequence from nucleotide 92 to nucleotide 1429 of SEQ ID NO:3 with at least one nucleotide substitution in the nucleotide sequence and wherein the taxol binding capacity of the beta-tubulin is not altered. Another exemplary DNA segment comprises the nucleotide sequence from nucleotide 92 to nucleotide 1429 of SEQ ID NO:3 with at least one nucleotide substitution in the nucleotide sequence and wherein the taxol binding capacity of the beta-tubulin is altered.

In another aspect, the invention is an amino acid sequence comprising at least a portion of a beta-tubulin of the fungal species *Pythium ultimum*. Preferably, the amino acid sequence comprises at least one taxol binding site. For some uses, it is preferable that the amino acid sequence is a protein having taxol binding site I and taxol binding site II. For amino acid sequences that can function as beta-tubulins, the amino acid sequence has taxol

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binding site I and taxol binding site II and is able to interact with alpha-tubulin. An exemplary amino acid sequence comprises at least a portion of the beta-tubulin as depicted in SEQ ID NO:4. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:4 comprising Amino Acids 1-31, with or without substitution. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:4 comprising Amino Acids 212-230, with or without substitution Another exemplary amino acid sequence comprises a portion of SEQ ID NO:4 comprising Amino Acids 212-230, wherein the amino acid at Amino Acid 219 is substituted. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:4 consisting essentially of Amino Acids 1-446 and wherein the portion behaves as a taxol-sensitive beta-tubulin. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:4 consisting essentially of Amino Acids 1-446 having at least one amino acid substitution that alters the taxol binding property of the portion. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:4 consisting essentially of Amino Acids 1-446 having at least one amino acid substitution that does not alter the taxol binding property of the portion. Another exemplary amino acid sequence is substituted with any amino acid which perturbs the three-dimensional structure of the amino acid sequence surrounding Amino Acid 219 as numbered in SEQ ID NO:4.

In another aspect, the invention is a purified DNA segment encoding a beta-tubulin of the fungal species *Phytophthora cinnamomi* or a portion thereof, wherein the DNA segment consists essentially of at least a portion of SEQ ID NO:5. An exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising the nucleotide sequence from nucleotide 11 through nucleotide 103. Another exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising the nucleotide sequence from nucleotide 11 through nucleotide 103, wherein at least one nucleotide in the nucleotide sequence is substituted, providing that when nucleotide substitution changes only one amino acid code nucleotide 80 cannot consist of adenine while nucleotide 81 is thymine and nucleotide 82 is adenine, cytosine or thymine. Another exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising nucleotide sequence from nucleotide 644 through nucleotide 700. Another exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising nucleotide sequence from nucleotide 700, wherein at least one nucleotide in the nucleotide sequence is substituted, providing that when nucleotide substitution changes only one amino acid code nucleotide 665 cannot be adenine while

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nucleotide 666 is adenine and nucleotide 667 is cytosine or thymine. Another exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising the nucleotide sequence from nucleotide 11 to nucleotide 1342 and wherein the DNA segment encodes a betatubulin. Another exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising the nucleotide sequence from nucleotide 11 to nucleotide 1342, wherein at least one nucleotide in the nucleotide sequence is substituted, providing that when nucleotide substitution changes only one amino acid code, nucleotide 665 cannot be adenine while nucleotide 666 is adenine and nucleotide 667 is cytosine or thymine. Another exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising the nucleotide sequence from nucleotide 11 to nucleotide 1342, wherein at least one nucleotide in the nucleotide sequence is substituted, providing that when nucleotide substitution changes only one amino acid code, nucleotide 665 cannot be adenine while nucleotide 666 is adenine and nucleotide 667 is cytosine or thymine, and wherein the taxol binding capacity of the beta-tubulin is not altered. Another exemplary DNA segment comprises a portion of SEQ ID NO:5 comprising the nucleotide sequence from nucleotide 11 to nucleotide 1342, wherein at least one nucleotide in the nucleotide sequence is substituted, providing that when nucleotide substitution changes only one amino acid code, nucleotide 665 cannot be adenine while nucleotide 666 is adenine and nucleotide 667 is cytosine or thymine, and wherein the taxol binding capacity of the beta-tubulin is altered.

In another aspect, the invention is an amino acid sequence comprising at least a portion of a beta-tubulin of the fungal species *Phytophthora cinnamomi* as depicted in SEQ ID NO:6. An exemplary amino acid sequence comprises a portion of SEQ ID NO:6 comprising Amino Acids 1-31. An exemplary amino acid sequence comprises a portion of SEQ ID NO:6 comprising Amino Acids 1-31, having at least one amino acid substituted, providing that when only one amino acid is substituted Amino Acid 24 is not isoleucine. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:6 comprising Amino Acids 212-230. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:6 comprising Amino Acids 212-230, having at least one amino acid substituted, providing that when only one amino acid is substituted Amino Acid 219 is not asparagine. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:6 comprising Amino Acids 212-230 with an amino acid substitution at Amino Acid 219, wherein the Amino Acid 219 is not substituted with asparagine. Another exemplary amino acid

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sequence comprises a portion of SEQ ID NO:6 consisting essentially of Amino Acids 1-446, wherein the portion behaves as a taxol-sensitive beta-tubulin. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:6 consisting essentially of Amino Acids 1-446, wherein the portion contains at least one amino acid substitution, providing that when only one amino acid is substituted Amino Acid 219 is not asparagine, and wherein the amino acid substitution alters the taxol binding property of the portion. Another exemplary amino acid sequence comprises a portion of SEQ ID NO:6 consisting essentially of Amino Acids 1-446, wherein the portion contains at least one amino acid substitution, providing that when only one amino acid is substituted Amino Acid 219 is not asparagine, and wherein the amino acid substitution does not alter the taxol binding property of the portion. Another exemplary amino acid sequence is substituted at Amino Acid 219 with any amino acid except asparagine which perturbs the three-dimensional structure of the amino acid sequence surrounding Amino Acid 219.

In another aspect, the invention is a vector comprising a purified DNA segment encoding a beta-tubulin of the fungal species *Pestalotiopsis microspora* or a portion thereof. Preferably, the vector comprises a portion encoding at least one taxol binding site.

In another aspect, the invention is a vector comprising a purified DNA segment encoding a beta-tubulin of the fungal species *Pythium ultimum* or a portion thereof. Preferably, the vector comprises a portion encoding at least one taxol binding site.

In another aspect, the invention is a vector comprising a purified DNA segment encoding a beta-tubulin of the fungal species *Phytophthora cinnamomi* wherein the DNA segment consists essentially of SEQ ID NO:5 or a portion thereof. Preferably, the vector comprises a portion encoding at least one taxol binding site.

In another aspect, the invention is a method of determining the taxol binding capacity of a beta-tubulin or beta-tubulin-like compound comprising providing antibodies raised against amino acid sequences comprising at least one taxol binding site of a beta-tubulin from a taxol-resistant *Pestalotiopsis microspora*, a taxol-sensitive *Pythium ultimum*, or taxol-sensitive *Phytophthora cinnamomi* as depicted in SEQ ID NO:6 to form a reagent, such antibodies distinguishing between taxol-binding and non-taxol-binding properties; contacting the beta-tubulin or beta-tubulin-like compound with the reagent; and determining

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degree of binding between the antibodies in the reagent and the beta-tubulin or beta-tubulinlike compound; whereby binding of antibodies raised against a taxol-resistant Pestalotiopsis microspora to the beta-tubulin or beta-tubulin-like compound indicates taxol resistance and whereby binding of antibodies which specifically recognize taxol-binding properties indicate taxol sensitive; whereby binding of antibodies which specifically recognize taxol-nonbinding properties indicate taxol resistance. In one embodiment, the antibodies in the reagent are raised against an amino acid sequence comprising at least one taxol binding site of a beta-tubulin from a taxol-resistant Pestalotiopsis microspora. In another embodiment, the antibodies in the reagent are raised against an amino acid sequence comprises at least one taxol binding site from SEQ ID NO:2. In another embodiment, the antibodies in the reagent are raised against an amino acid sequence comprising at least one taxol binding site of a beta-tubulin from a taxol-sensitive Phythium ultimum. In another embodiment, the antibodies in the reagent are raised against an amino acid sequence comprises at least one taxol binding site from SEQ ID NO:4. In another embodiment, the antibodies in the reagent are raised against an amino acid sequence comprising at least one taxol binding site of a beta-tubulin from a taxol-sensitive Phytophthora cinnamomi as depicted in SEQ ID NO:6. In another embodiment, the antibodies in the reagent are raised against an amino acid sequence comprises at least one taxol binding site from SEQ ID NO:678. In this method, the beta-tubulin or beta-tubulin-like compound are selected from the group consisting of recombinantly expressed protein, exogenously isolated protein, synthetic peptides, and cell cultures.

In another aspect, the invention is a method of screening a composition of matter for the presence of taxol or taxol-like compounds comprising providing beta-tubulins with amino acid sequences comprising both taxol binding sites from *Pythium ultimum* or taxol-sensitive *Phytophthora cinnamomi* as depicted in SEQ ID NO:6 in addition to alpha-tubulin from any taxol-sensitive organism to form a reagent; contacting the composition of matter with the reagent; and determining the ability of the composition of matter to promote MT assembly or ability to prevent depolymerization of assembled MTs under depolymerizing conditions; whereby the ability to promote microtubule assembly or prevent depolymerization indicate the possible presence of taxol or taxol-like compounds in the composition of matter.

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In another aspect, the invention is a method of screening a composition of matter for the presence of taxol or taxol-like compounds comprising providing mycelia of taxol-sensitive *Pythium ultimum* or a taxol-sensitive *Phytophthora cinnamomi* which harbors betatubulin in SEQ ID NO:6; contacting the composition of matter with the mycelia in the presence of the labeled taxol; and determining the degree of competitive inhibition of binding between the beta-tubulins and the labeled taxol by the composition of matter, whereby the composition of matter is determined to possess taxol or taxol-like compounds if it is able to block labeled taxol binding to the beta-tubulins from the taxol-sensitive *Pythium ultimum* or *Phytophthora cinnamomi*.

In another aspect, the invention is a method of altering the taxol binding property of a recombinantly expressed beta-tubulin or a portion thereof comprising determining the identity of the codon at Amino Acid 219 as numbered in SEQ ID NO:1 in the coding region of the vector; and if the codon at Amino Acid 219 codes for any amino acid except threonine, substituting nucleotides in the codon to code for threonine at Amino Acid 219 to alter a non-taxol-binding beta-tubulin or portion thereof to a taxol-binding beta-tubulin or portion thereof, or if the codon at Amino Acid 219 codes for threonine, substituting nucleotides in the codon to code for any amino acid except threonine at Amino Acid 219 to alter a taxol-binding beta-tubulin or portion thereof to a non-taxol-binding beta-tubulin or portion thereof.

In another aspect, the invention is a method of developing a taxol-sensitive fungal cell from a taxol-resistant fungal cell comprising transforming the non-taxol-sensitive fungal cell by introducing a DNA segment encoding taxol-binding beta-tubulin comprising threonine at Amino Acid 219 as numbered in SEQ ID NO:2; wherein the transformed fungal cell expresses the DNA segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.

In another aspect, the invention is a transgenic taxol-sensitive fungal cell transformed by introducing a DNA segment encoding taxol-binding beta-tubulin comprising threonine at Amino Acid 219 as numbered in SEQ ID NO:2, wherein the transformed fungal cell expresses the DNA segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.

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In another aspect, the invention is a method of developing a taxol-resistant fungal cell from a taxol-sensitive fungal cell comprising transforming the taxol-sensitive fungal cell by introducing a DNA segment encoding non-taxol-binding beta-tubulin wherein the amino acid at Amino Acid 219 as numbered in SEQ ID NO:2 is not threonine; wherein the transformed fungal cell over-expresses the DNA segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.

In another aspect, the invention is a transgenic taxol-sensitive fungal cell transformed by introducing a DNA segment encoding taxol-binding beta-tubulin comprising threonine at Amino Acid 219 as numbered in SEQ ID NO:2, wherein the transformed fungal cell over-expresses the DNA segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.

In another aspect, the invention is a method of screening a composition of matter for the presence of taxol or taxol-like compounds comprising providing distinguishable taxol-resistant and taxol-sensitive fungal cells; contacting the composition of matter with the fungal cells; and determining the growth of inhibition of the fungal cells; whereby the composition of matter is determined to possess taxol or taxol-like compounds if it is able to inhibit the growth of taxol-sensitive fungal cells but not able to inhibit the growth of taxol-resistant fungal cells. The method can be performed wherein the distinguishable taxol-resistant and taxol-sensitive fungal cells consists essentially of transgenic taxol-resistant and taxol-sensitive isogenic fungal cells. The method can also be performed with taxol-resistant fungal cells derived from one fungus which is unrelated to the fungi from which the taxol-sensitive fungal cells are derived.

In another aspect, the invention is a method for controlling the growth of a plant pathogen comprising determining the taxol sensitivity of the plant pathogen; and if the pathogen is determined to be taxol-sensitive, the plant and soil surrounding the plant are treated with a taxol-producing *P. microspora*. In an exemplary method, the taxol sensitivity of the plant pathogen is determined by identifying Amino Acid 219, wherein the plant is designated as taxol-sensitive if Amino Acid 219 is threonine.

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BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 is a graph depicting the effect of taxol on mycelial growth in *P. microspora*, *P. ultimum*, *P. cinnamomi* and *A. klebsiana*. Fungal mycelia were grown on potato dextrose agar (PDA) plates containing different concentrations of taxol. The inhibitory effect of taxol was assessed by colony diameter, and compared to mycelia grown in the absence of taxol. Experiments were conducted in duplicate, and data presented are an average of several experiments.

Fig. 2 depicts the nucleotide and deduced amino acid sequence of beta-tubulin from *P. microspora* Ne32 cDNA, TUBB-pm. Numerals on the left indicate nucleotide position, and numerals on the right indicate amino acid position. The sequences of the gene-specific primers NETUB5 and NETUB6 are underlined. The translation initiation codon ATG is underlined, the translation termination codon is marked by an asterisk, and the putative polyadenylation signal is double underlined.

Fig. 3 depicts the nucleotide and deduced amino acid sequence of beta-tubulin from *P. ultimum* cDNA, TUBB-pu. Numerals on the left indicate nucleotide position, and numerals on the right indicate amino acid position. The sequences of the gene-specific primers WT1L-U and WT1L-L are underlined. The translation initiation codon ATG is underlined, the translation termination codon is marked by an asterisk, and the two putative polyadenylation signals are double underlined. The arrow at nucleotide 1507 indicates the position of the poly (A) tract in the shorter 1537 bp cDNA.

Fig. 4 depicts the nucleotide and deduced amino acid sequence of beta-tubulin from *P. cinnamomi* cDNA, TUBB-pc. Numerals on the left indicate nucleotide position, and numerals on the right indicate amino acid position. The sequences of the gene-specific primers PCBTUB1U, PCBTUB2U and PCBTUB4L are underlined. The translation initiation codon ATG is marked by ###, and the translation termination codon is marked by an asterisk. Nucleotides and amino acids which differ between TUBB-pc and the sequence U22050 (Genbank accession number) reported by Weerakoon et al. (Weerakoon et al. 1998. "Isolation and characterization of the single β-tubulin gene in *Phytophthora cinnamomi*," *Mycologia* 90:85-95) are double underlined.

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Fig. 5 compares the amino acid sequence of *P. cinnamomi* beta-tubulins ("TUBB-pc") reported herein and previously by Weerakoon et al. (Weerakoon, et al. 1998. *Mycologia* 90:85-95) ("U22050"). The deduced amino acid sequence of *P. cinnamomi* TUBB-pc is shown in its entirety, and the eight residues (Amino Acids 24, 219, 249, 251-253, 359, and 428) which differ from the U22050 sequence (Genbank accession number) reported by Weerakoon et al. (1998) are shown below the TUBB-pc sequence. Amino Acids 1-31 and 212-231 (denoted herein as taxol binding region I and II, respectively) are indicated by a line above the sequence.

Fig. 6A and 6B depict the amino acid sequence alignment of beta-tubulins. The alignment was obtained using the ClustalW alignment program. The amino acid sequence of *P. microspora* beta-tubulin is shown in its entirety, and residues which differ in other beta-tubulins are shown below. Numerals on the right indicate amino acid positions. Sequences underlined indicate regions important for GTP binding (Amino Acids 63-77), phosphate binding (Amino Acids 140-146), and Mg²⁺ binding (Amino Acids 203-206). Amino Acids 1-31 and 212-231 (denoted here as taxol binding region I and II, respectively) are indicated by a line above the sequence. Amino Acids Phe270, Leu273 and Ser364 are marked above with #. Amino acids which are important for fungal resistance to benzimidazoles (Amino Acids 6, 165, 167, 198, 200 and 241) are marked above by asterisks. Gaps in alignment are indicated by dashes, and the end of each sequence is marked by "\$". The Genbank accession numbers for beta-tubulins from *N. crassa*, *A. midulans* benA, *A. klebsiana* and human β2 are listed in Table I. The *P. cinnamomi* depicted is SEQ ID NO:6.

Fig. 7A and 7B are graphs depicting the specific binding of [³H]taxol to *P. ultimum* but not to *P. microspora*. Fig. 7A demonstrates that specific binding of [³H]taxol to *P. ultimum* increased as a function of [³H]taxol concentration, while *P. microspora* showed no or very little specific binding. Actively growing fungal cells were incubated with different concentrations of [³H]taxol at room temperature for 2 hours before quenching. Specific binding was calculated as the difference between binding of [³H]taxol in the presence and absence of a 100-fold excess of unlabeled taxol. Specific binding represents 30-70% of the total binding to *P. ultimum* but less than 5% of the total binding to *P. microspora*. Binding of [³H]taxol to *P. microspora* cells was performed in the presence of Triton X-100 (0.1%)

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v/v) to disrupt the cell membrane. Fig. 7B depicts the reduction of specific binding of [³H]taxol to *P. ultimum* in the presence and absence of the microtubule-depolymerizing drug thiabendazole in a dose-dependent manner. Cells were either not treated (0 μM) or treated with different concentrations of thiabendazole for three hours at room temperature.

Subsequently, cells were incubated with 5 nM [³H]taxol in the presence or absence of a 100-fold excess of unlabeled taxol for two hours before quenching. The specific binding of [³H]taxol to untreated cells was defined as 100%. The experiments depicted in Fig. 7A and 7B were conducted in duplicate, and data presented are representative of several experiments.

Fig. 8 depicts the amino acid sequences of the taxol binding region I (Amino Acids 1-31) and II (Amino Acids 212-231) of beta-tubulins from different organisms. The amino acid sequences of the taxol binding regions I and II for pig beta-tubulin are shown in their entirety and residues which differ are shown for other beta-tubulins. The taxol sensitivity of each organism is indicated, "s" for sensitive and "r" for resistant. Amino Acids 15-25 and 212-222, which have been shown to be involved in taxol binding by both cross-linking and electron crystallography, are marked with asterisks. The taxol binding region II of A. klebsiana is between Amino Acids 211-230 due to a gap in its sequence. Pig beta-tubulin is described by Nogales, et al. (Nogales, et al. 1999. Nature 391:199-203), and Genbank accession numbers for other sequences are listed in Table I. The sequence for P. cinnamomi presented herein is depicted in SEQ ID NO:6.

DETAILED DESCRIPTION

One aspect of the present invention is an isolated gene comprising an open reading frame coding for the protein beta-tubulin or a portion thereof. The corresponding cDNA have been isolated and characterized for taxol-resistant *Pestalotiopsis microspora* Ne32, taxol-sensitive *Pythium ultimum*, and taxol-sensitive *Pythium cinnamomi*. The nucleotide and deduced amino acid sequences of beta-tubulin for *Pestalotiopsis microspora* Ne32 are given in SEQ ID NO:1 and SEQ ID NO:2, respectively; for *Pythium ultimum*, in SEQ ID NO:3 and SEQ ID NO:4, respectively; and for *Pythium cinnamomi*, in SEQ ID NO:5 and SEQ ID NO:6, respectively. Through characterization of the taxol sensitivity of the beta-tubulins encoded by the genes of the present invention, it has been found that the identity of Amino Acid 219 of beta-tubulin as numbered in SEQ ID NO:2, SEQ ID NO:4, or SEQ ID

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NO:6 is an indicator of taxol sensitivity. The presence of threonine at Amino Acid 219 ("Thr219") indicates taxol sensitivity, while the presence of asparagine ("Asn219") or glutamine ("Gln219") indicate taxol resistance.

In another aspect, the present invention is the beta-tubulin protein or protein fragments encoded by the novel genes disclosed herein. Since the *P. ultimum* and *P. cinnamomi* beta-tubulin proteins of the present invention are capable of binding taxol, proteins and protein fragments comprising taxol-binding sites derived from the genes coding for beta-tubulin described herein can be produced by heterologous expression in *E. coli* and other systems, purified by standard procedures, and used in an *in vitro* assay for detecting taxol and taxol-like substances by using methods well known in the art (Schiff, et al. 1979. *Nature* 277:665-667). For example, the beta-tubulin proteins of the present invention can be used to screen plant or fungal extracts as well as synthetic compounds for taxol or taxol-like substances as possible anticancer drugs. Beta-tubulins produced by making specific amino acid substitutions, deletions, or alterations can be used as experimental tools to further determine the molecular basis of taxol binding to the beta-tubulin protein.

In another aspect, antibodies (polyclonal or monoclonal) raised against all or portions of the beta tubulins of the present invention can be used to determine if a composition of matter has taxol binding properties. In one method, antibodies capable of binding to taxol-sensitive beta-tubulin and/or taxol-resistant beta tubulins are exposed to a composition of matter prepared for in situ hybridization (Ausubel, et al. 1997. Current Protocols in Molecular Biology, John Wiley & Sons), Elisa, or Western blot. Visualization of antibody-antigen binding is mediated through any means known in the art, e.g., secondary radiolabeled or fluorescent antibodies or colorimetric methods using peroxidase and/or alkaline phosphatase (Harlow, E.D. and Lane, D. 1988. Antibodies: A Laboratory Manual). For example, antibodies raised to a portion of SEQ ID NO:4 comprising Amino Acid 219 would bind to a beta-tubulin which had threonine at Amino Acid 219 but would not bind to a beta-tubulin having a different amino acid at Amino Acid 219, so that detectable binding would indicate the presence of threonine at Amino Acid 219, and hence, sensitivity to taxol. This type of assay is useful for screening a variety of compositions of matter, including living matter such as plants or microorganisms, or non-living matter such as plant materials, patient samples, or compound libraries for the presence of beta-tubulin.

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In another aspect, the present invention is a method of designing taxol analogs or other compounds which mimic the interaction of taxol with beta-tubulin based on the identification of specific amino acids in the beta-tubulins corresponding to taxol-binding and taxol-sensitivity. The previously reported three-dimensional structure (Nogales, et al. 1998. *Nature* 391:199-203) can be applied to developing and optimizing antineoplastic and antifungal compounds with respect to Amino Acid 219 and the surrounding area. Further, such information can also be used to generate mutant beta-tubulins with altered taxol sensitivity by substituting amino acids at specific positions in the beta-tubulin protein.

In another aspect, the present invention is a method of generating isogenic strains of fungi using a gene of the present invention, which allows studies of taxol related pharmacology to be performed against a known background. Further, the present invention is a method of using these isogenic fungal strains, one of which is taxol sensitive and the other taxol resistant, to screen plant extracts, fungal extracts, extracts from other organisms, and synthetic compounds for taxol-like substances as possible anticancer agents. The present invention is also a method of using two unrelated fungal strains, one of which is taxol sensitive and the other taxol resistant, to screen plant extracts, fungal extracts, extracts from other organisms, and synthetic compounds for taxol-like substances as possible anticancer agents.

The genes and proteins of the present invention are characterized in the following examples. It is to be understood that the examples are exemplary of the invention and are intended to be illustrative of the invention, but are not to be construed to limit the scope of the invention in any way. Modifications may be made in the structural features of the invention without departing from the scope of the invention.

Example 1: Differential taxol sensitivity in selected fungi

Taxol sensitivity was established for the fungal strains used in the isolation of the beta-tubulin cDNAs of the present invention.

Pestalotiopsis microspora strain Ne32, previously disclosed in U.S. Patent No. 5,861,302, was licensed from Montana State University. Pythium ultimum (ATCC 26083), Achlya klebsiana (ATCC 52605), and Pythium cinnamomi (ATCC 200982) were purchased

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from American Type Culture Collection (Manassas, VA). Taxol was obtained from Sigma Chemical Company (St. Louis, MO).

The effect of taxol on the growth of *P. microspora* Ne32, *P. ultimum*, and *P. cinnamomi* was examined. For comparison, *A. klebsiana*, an oomycete closely related to *P. ultimum*, was also included in these experiments. As shown in Fig. 1, the growth of *P. microspora* was highly resistant to taxol up to 11.7 μM. By comparison, *A. klebsiana* showed moderate sensitivity, and its growth was reduced by 40% in 11.7 μM taxol. Finally, *P. ultimum* and *P. cinnamomi* were shown to be the most sensitive of the four species. Growth of *P. ultimum* and *P. cinnamomi* was inhibited even at low concentrations of taxol (IC₅₀ 0.1 μM). This sensitivity is comparable to the level of taxol (0.25 μM) that inhibits Hela cell division (Schiff, et al. 1979. *Nature* 277:665-667).

Example 2: Isolation of β -tubulin cDNA sequences

Beta-tubulin cDNA sequences were determined for *P. microspora* Ne32, *P. ultimum*, and *P. cinnamomi* from RNA isolated from fungal mycelia. Automated dideoxynucleotide sequencing was performed by a contracting laboratory. Sequence comparison was performed using the BLAST program at the Internet site of the National Center for Biotechnology Information. The amino acid sequence alignment was performed using ClustalW program, and other analysis using MacVector program.

To isolate beta-tubulin cDNA sequences from *P. microspora* Ne32 and *P. ultimum*, four degenerate primers were designed according to conserved motifs in fungal beta-tubulin amino acid sequences. A forward degenerate primer BTUB1, 5'-CTGGGCYAAGGGYC AYTACACYGAG-3' (SEQ ID NO.7, was designed corresponding to amino acid residues Trp-Ala-Lys-Gly-His-Tyr-Thr-Glu (or WAKGHYTE in single letter amino acid code; SEQ ID NO:8); a reverse primer BTUB2, 5'-CGAAGAARTGRARNCGRGGGAARGG-3' (SEQ ID NO:9), corresponding to amino acid residues Pro-Phe-Pro-Arg-Leu-His-Phe-Phe (or PFPRLHFF in single letter amino acid code; SEQ ID NO:10); a forward primer BTUB3, 5'-CGAGCCYTACAACGCYACYCT-3' (SEQ ID NO:11), corresponding to amino acid residues Glu-Pro-Tyr-Asn-Ala-Thr-Leu (or EPYNATL in single letter amino acid code; SEQ ID NO:12); and a reverse primer BTUB4, 5'-CTCGTTCATGTTRSWCTCRGCCTC-3' (SEQ ID NO:13), corresponding to amino acid residues Glu-Ala-Glu-Ser-Asn-Met-Asn-Asp

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(or EAESNMND in single letter amino acid code; SEQ ID NO:14). All primers were synthesized by a contracting laboratory according to our specifications.

In order to isolate beta-tubulin cDNA from P. microspora Ne32, five micrograms (5 ug) of total RNA from mycelia grown for 5 days was used to synthesize first-strand cDNA using primer BTUB4 in a 20 microliter (20 µl) reaction. One microliter (1 µl) of the cDNA product was used as template in Polymerase Chain Reactions (PCR). The cycling program contained 8 cycles of an annealing temperature of 52°C, followed by 22 cycles with an annealing temperature of 62°C. Degenerate primers BTUB3 and BTUB4 generated an amplification product of 0.8 kb, whereas BTUB2 and BTUB3 generated a product of 0.3 kb. The desired bands were gel-purified using Geneclean (BIO 101, Vista, CA), ligated into the pPCR2.1 vector (Invitrogen; San Diego, CA), and transformed into E. coli XL1 -Blue cells. Inserts were sequenced and used to synthesize a gene-specific forward primer NETUB5, 5'-GGGTGTCACCACTTGCTTGCGTTT-3' (SEQ ID NO:15), and a reverse primer NETUB6, 5'-TCGAGTTTCCGACGAAAGTGGACGA-3' (SEQ ID NO:16). To obtain full-length clones, a Marathon cDNA Library was constructed using one microgram (1 µg) mRNA according to manufacturer (Clontech; Palo Alto, CA). One microliter (1 µl) of this library was diluted 250-fold, and five microliters (5 µl) were used in Rapid Amplification of cDNA Ends (RACE) reactions using the PCR cycling program recommended by the manufacturer. For 5' RACE, library adaptor primer AP1 (Clontech) and primer NETUB6 generated a product of 1.3 kb. For 3' RACE, primers AP1 and NETUB5 generated a product of 1.0 kb. The desired bands were gel-purified and cloned into the pPCRII-TOPO vector (Invitrogen, Carlsbad, CA). The region between 1 to 1105 bp from the 5' RACE product was ligated at an internal BamHI site with the region between 1106 to 1668 bp from the 3' RACE product to form the composite cDNA (Fig. 2). The resulting composite cDNA from P. microspora was 1668 bp long, designated as TUBB-pm, and its nucleotide and deduced amino acid sequence are shown in SEQ ID NO:1 and SEQ ID NO:2, respectively, as well as Fig 2. This cDNA encodes a protein of 446 amino acids with a calculated Mr of 49,832 and pI of 4.6. It contains 74 nucleotides in the 5' untranslated region (UTR), and 229 nucleotides in the 3' UTR followed by a 24 nucleotide poly (A) tail. A sequence AATAA (nucleotides 1539-1543 of SEQ ID NO:1) with the closest similarity to the animal and viral polyadenylation signal AATAAA (Proudfoot, N.J. and Brownlee, G.G. 1976. "3' Non-coding region

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sequences in eukaryotic messenger RNA," *Nature* 263:211-214) was located 103 bp upstream of the poly (A) tract.

In order to isolate beta-tubulin cDNA from P. ultimum, five micrograms (5 µg) of total RNA from mycelia grown for six days was used to synthesize first strand cDNA with oligo-dT primer (GibcoBRL; Gaithersburg, MD) in a twenty microliter (20 µl) reaction. Two microliters (2 µl) of cDNA product were used as the template in PCR reactions with a cycling program similar to that described above. Degenerate primers BTUB1 and BTUB4 generated a product of 1.0 kb, whereas BTUB1 and BTUB2 amplified a product of 0.5 kb. The desired bands were gel-purified and ligated into the pPCR2.1 vector. Inserts were sequenced and used to design a gene-specific forward primer WT1L-U, 5'-CTAT CATGTGCACGTACTCGGTGTGC-3' (SEQ ID NO:17), and a reverse primer WT1L-L, 5'-CTGGGACGGTCAAAGCACGGTACTGC-3' (SEQ ID NO:18). For 5' RACE, first-strand cDNA was synthesized using primer WT1L-L and used as template in PCR reactions. Primers WT1L-L and Cap-Switch (Clontech) generated a 0.95 kb product using Advantage-GC cDNA PCR kit (Clontech). For 3' RACE, first strand cDNA was synthesized using CapFinder cDNA Library Construction kit (Clontech). With the resulting cDNAs as template, primers WTIL-U and CDS/3' (Clontech) generated two PCR products of 1.0 and 1.1 kb, respectively. PCR fragments were gel-purified and cloned into the pPCR2.1 vector. In P. ultimum, isolated tubulin cDNAs were of two types, one composite cDNA was 1650 bp long, and the other was 1537 bp long. These two cDNAs differ only in the position at which the poly (A) tail has been added. The region between 1 to 824 bp from the 5' RACE product was ligated at an internal MfeI site with the region between 825 to 1650 bp from the 1.1 kb 3' RACE product to form the composite 1650 bp cDNA designated as TUBB-pu, and its nucleotide and deduced amino acid sequence are shown in SEQ ID NO:3 and SEQ ID NO:4, respectively, as well as Fig. 3. This cDNA encodes a protein of 446 amino acids with a calculated Mr of 50,047 and pI of 4.6. It contains 91 nucleotides in the 5' UTR, and 199 nucleotides in the 3' UTR followed by a 19 nucleotide poly (A) tract. Two imperfect polyadenylation signals were tentatively identified, ATATA at 57 bp upstream of poly (A) tract in the 1537 bp cDNA (nucleotides 1445-1449 of SEQ ID NO:3), and AATATT at 80 bp upstream of poly (A) tail in the 1650 bp cDNA (nucleotides 1546-1551 of SEQ ID NO:3). The sizes of these cDNAs match well with transcript sizes in Northern analysis as shown below, indicating they are complete.

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Four gene-specific primers were synthesized based on the reported beta-tubulin cDNA sequence from *P. cinnamomi* (Weerakoon, et al. 1998. *Mycologia* 90:85-95). The forward primer PCBTUB1U (5'-CAGCGACAACATGAGAGAGCTCG-3'; SEQ ID NO:19) corresponds to region 270-292 in its cDNA sequence, the forward primer PCBTUB2U (5'-CGATGAGGTCATGTGCCTGGATAA-3'; SEQ ID NO:20) corresponds to region 867-890; the reverse primer PCBTUB3L (5'-AAACGGAGGCACGTGGTGATG-3'; SEQ ID NO:21) corresponds to region 984-1005; the reverse primer PCBTUB4L (5'-CGCGTC TATCTCATCCATTCCTCG-3'; SEQ ID NO:22) corresponds to region 1596-1619.

Five micrograms (5 µg) of total RNA from P. cinnamomi mycelia grown for 5 days was used to synthesize first-strand cDNA using oligo-dT primer in a 20 µl reaction. One microliter (1 µl) of the cDNA product was used as template for PCR. The cycling program comprised 30 cycles with an annealing temperature of 62°C. Primer PCBTUB1U and PCBTUB4L generated an amplification product of 1.3 kb, and primer PCBTUB2U and PCBTUB4L generated an amplification product of 0.75 kb. Primer PCBTUB1U or PCBTUB2U in combination with PCBTUB3L did not generate any product. The desired bands were gel-purified using Geneclean (BIO101), ligated into the pPCR2.1 vector (Invitrogen), and transformed into E. coli XL1-Blue cells. Four clones of the 1.3 kb fragment (clone # C16-1, 4, 9 and 10) and one clone of the 0. 75 kb fragment (clone # C10-5) were sequenced from both directions and found to conform to the same sequence. The nucleotide and deduced amino acid sequence of the 1.3 kb long cDNA, designated as TUBB-pc, are shown in SEQ ID NO:5 and SEQ ID NO:6, respectively, as well as Fig. 4. It encodes a 444 amino acid long beta-tubulin protein, with a calculated Mr of 50 kDa, and a pI of 4.7. There are 10 nucleotides in the 5' untranslated region (UTR), and 5 nucleotides in the 3' UTR.

A sequence encoding *P. cinnamomi* beta-tubulin has been previously reported (Weerakoon, et al. 1998. *Mycologia* 90:85-95; Genbank accession number U22050), and the deduced amino acid sequence of *P. cinnamomi* beta-tubulin disclosed herein was compared to that disclosed by Weerakoon et al. The TUBB-pc cDNA sequence shown in SEQ ID NO:5 and Fig. 4 differs by 36 nucleotides (2.7%) within the coding region from the one reported by Weerakoon et al. An alignment of the beta-tubulin amino acid sequences deduced from TUBB-pc (SEQ ID NO:6) and the one previously reported by Weerakoon

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("U22050"; SEQ ID NO:23) is shown in Fig 5. The two sequences differ by 8 amino acids. Four are conserved changes, while the other four are nonconserved changes. One change is within each of the taxol-binding sites. In the taxol-binding region I (Amino Acids 1-31), Amino Acid V24 (valine at Amino Acid 24) in TUBB-pc differs from I24 (isoleucine at Amino Acid 24) in U22050. In the taxol-binding region II (Amino Acids 212-231), Amino Acid T219 (threonine at Amino Acid 219) in TUBB-pc differs from N219 (asparagine at Amino Acid 219) in U22050.

The deduced amino acid sequence of beta-tubulin from P. microspora (SEQ ID NO:2), P. ultimum (SEQ ID NO:4), and P. cinnamomi (SEQ ID NO:6) show features expected of beta-tubulin, as shown by an alignment with human \(\beta\)2-tubulin (SEQ ID NO:24) and from beta-tubulins from Neurospora crassa (SEQ ID NO:25), A. nidulans benA (SEQ ID NO:26), and A. klebsiana (SEQ ID NO: 27) depicted in Fig. 6. These sequences can be divided into N-terminal (Amino Acids 1-205), intermediate (Amino Acids 206-381) and Cterminal domains (Nogales, et al. 1998. Nature 391:199-203). Their N-terminal domain contains conserved motifs important for GTP binding [Ala-Ile-Leu-Val-Asp-Leu-Glu-Pro-Gly-Thr-Met-Asp-Ser-Val-Arg or AILVDLEPGTMDSVR in single letter amino acid code (SEQ ID NO:28) and Ala-Val-Leu-Val-Asp-Leu-Glu-Pro-Gly-Thr-Met-Asp-Ser-Val-Arg or AVLVDLEPGTMDSVR in single letter amino acid code (SEQ ID NO:29) between Amino Acids 63-77 in SEQ ID NO:2, SEQ ID NO:4, SEQ ID NO:6, SEQ ID NO:24, SEQ ID NO:25, and SEQ ID NO:26, and between Amino Acids 62-76 in SEQ ID NO:27], phosphate binding [Gly-Gly-Gly-Thr-Gly-Ser-Gly or GGGTGSG in single letter amino acid code (SEQ ID NO:30) between Amino Acids 140-146 in SEQ ID NO:2, SEQ ID NO:4, SEQ ID NO:6, SEQ ID NO:24, SEQ ID NO:25, and SEQ ID NO:26, and between Amino Acids 139 and 145 in SEQ ID NO:27], and Mg²⁺ binding (Asp-Asn-Glu-Ala or DNEA in single letter amino acid code (SEQ ID NO:31) between Amino Acids 203-206 in SEQ ID NO:2, SEQ ID NO:4, SEQ ID NO:6, SEQ ID NO:24, SEQ ID NO:25, and SEQ ID NO:26, and between Amino Acids 202-205 in SEQ ID NO:27) (Linse, K. and Mandelkow, E.M. 1988. "The GTP-binding peptide of β-tubulin: localization by direct photoaffinity labeling and comparison with nucleotide-binding proteins," J Biol Chem 263:15205-15210; and Farr, G.W. and Stemlicht, H. 1992. "Site-directed mutagenesis of the GTP-binding domain of \betatubulin," J Mol Biol 227:307-321). In addition, the N-terminal Met-Arg-Glu-Ile (or MREI

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in single letter amino acid code; (SEQ ID NO:32) motif has been shown to autoregulate the stability of beta-tubulin mRNA in animal cells (Yen, et al. 1988. "Autoregulated instability of β-tubulin mRNAs by recognition of the nascent amino terminus of β-tubulin," *Nature* 334:580-585). This motif and a variant Met-Arg-Glu-Leu (or MREL in single letter amino acid code; SEQ ID NO:33) are present in the fungal beta-tubulins shown in Fig. 6, and they may function similarly. The C-terminal domain has been reported to be important for interactions with microtubule-associated proteins (MAPS) and motor proteins (Nogales, et al. 1998 Nature 391:199-203). Sequences near the C-terminus are hypervariable and acidic, a common feature of beta-tubulins (Sullivan, K.F. 1988. "Structure and utilization of tubulin isotypes," *Ann Rev Cell Biol* 4:687-716).

The amino acid sequence of beta-tubulins from different organisms are well conserved and exhibit at least 63% identity (Oakley, B.R. 1994. "Gamma-tubulin." In: Hyams JS, Lloyd CW (eds) *Microtubules*. Wiley-Liss, New York, pp. 38-45). Table I shows the percentage identity between the beta-tubulin amino acid sequence of *P. microspora* and *P. ultimum* with beta-tubulins of other organisms. The beta-tubulin from *P. microspora* shares the highest identity (93-97%) with filamentous ascomycetes such as *A. flavus*, *A. nidulans* benA and *N. crassa*, and lower identity (73-78%) with single-cell ascomycetes and oomycetes. Its identity (78-85%) with beta-tubulin from non-fungal organisms is also relatively low. In contrast, beta-tubulin from *P. ultimum* shows the highest identity (96-97%) with beta-tubulin from two oomycetes, *A. klebsiana* and *P. cinnamomi*, but shares limited identity (71-78%) with beta-tubulin from ascomycetes. The beta-tubulin from *P. ultimum* also shows relatively high identity (86-93%) with beta-tubulin from non-fungal organisms such as the green algae *C. reinhardtii*, the protozoa *T. thermophila*, the slime mold *Physarum polycephalum*, and various animals.

Example 3: Extraction of genomic DNA and Southern analysis

Since multiplicity within the genes that encode beta-tubulin could affect the taxol-dependent property of microtubules, Southern analysis was used to determine whether *P. microspora* Ne32 and *P. ultimum* harbor one or more copies of beta-tubulin gene.

Mycelia of *P. microspora* Ne32 or *P. ultimum* grown for three days were harvested, and genomic DNA was isolated using Elu-Quik Hi-Volume Genomic kit (Schleicher

Table I: Amino Acid Sequence Identity Between Beta-tubulin from P. microspora or P. ultimum and Other Organisms^a

		Percentage of Identical Amino Acids		
Organism	Accession No.	P. Microspora	P. ultimum	
Saccharomyces cerevisiae	VO 1296	73	71	
Schizosaccharomyces pombe	M10347	78	76	
Aspergillus flavus	M38265	94	77	
Aspergillus nidulans benA	M17519	93	77	
Neurospora crassa	A25377	97	78	
Achlya klebsiana	P20802	78	97	
Phytophthora cinnamomi	U22050	77	96	
Chlamydomonus reinhardtii (β1 and β2)	P04690	78	91	
Tetrahymena thermophila (β1 and β2)	P41352	80	93	
Physarum polycephalum (slime mold) (β1)	P07436	81	90	
Drosophila melanogaster (fruit fly) (β1)	M20419	83	86	
Xenopus laevis (β4)	P30883	85	89	
chicken (β2)	P32882	84	88	
mouse (β3)	C25437	84	89	
human (β2)	P05217	84	89	

^a The amino acid sequences of beta-tubulin were retrieved from Genbank or Swiss-Prot. Pairwise identity was performed either with ClustalW or BLAST program.

Schuelle, Neene, NH). Five micrograms (5 µg) of genomic DNA was digested with restriction enzymes for 4 hours at 37°C, separated on 0.8% agarose gel, transferred to Nylon filters (Tropix, Bedford, MA), and wet filters were cross-linked using GS Gene Linker UV Chamber (Biorad; Hercules, CA). Southern blotting was performed under stringent conditions according to standard protocols (Sambrook et al. 1989. *Molecular Cloning: A*

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Laboratory Manual. Cold Spring Harbor Laboratory, Cold Spring Harbor, NY). A 413 bp PCR fragment of P. microspora beta-tubulin was generated with primers NETUB5 and NETUB6, while a 377 bp PCR fragment of P. ultimum beta-tubulin was generated with primers WT1L-U and WT1L-L. PCR products were gel-purified and labeled with α - ^{32}P dCTP using Ready-to-Go beads (Pharmacia, Piscataway, NJ) by random priming. Probes were purified with Micro Bio-Spin columns (Biorad).

In P. microspora, the beta-tubulin probe hybridized to a single band from genomic DNA digested with EcoRI, HindIII, or Sall, and two bands from BamHI digested sample. In P. ultimum, the beta-tubulin probe hybridized to a single band from genomic DNA digested with BamHI, SalI, PvuI, or PstI, and two bands from EcoRI digested sample. The sizes of these fragments match those predicted from the restriction endonuclease map of the corresponding cDNA clones. Since beta-tubulin genes are typically highly conserved, these results show that both P. microspora and P. ultimum contain a single copy of the betatubulin gene, consistent with previous reports that fungi generally have one, or at most a few, beta-tubulin genes (Neff, et al. 1983. "Isolation of the β-tubulin gene from yeast and demonstration of its essential function in vivo, "Cell 33:211-219; Hiraoka, et al. 1984. "The NDA3 gene of fission yeast encodes β -tubulin: a cold sensitive *nda3* mutation reversibly blocks spindle formation and chromosome movement in mitosis," Cell 39:349-358; Orbach, et al. 1986. "Cloning and characterization of the gene for \beta-tubulin from a benomylresistant mutant of Neurospora crassa and its use as a dominant selectable marker," Mol Cell Biol 6:2452-2461; Cameron et al. 1990. "Cloning and analysis of β-tubulin gene from a protoctist," JBiol Chem 265:15245-15252; and Weerakoon et al. 1998. Mycologia 90:85-95; May, et al. 1987. "Aspergillus nidulans β-tubulin genes are usually divergent," Gene 55:231-243; Panaccione et al. 1988. "Colletotrichum graminicola transformed with homologous and heterologous benomyl-resistance genes retains expected pathogenicity to corn," Mol Plant Microbe Interact 1:113-120; and Goldman et al. 1993. "A nucleotide substitution in one of the β -tubulin genes of *Trichoderma viride* confers resistance to the antimitotic drug methyl benzimidazole-2-yl-carbamate," Mol Gen Genet 240:73-80). The fact that P. microspora and P. ultimum have only a single copy of the beta-tubulin gene indicates that it is responsible for the sensitive or resistance properties of the organism to taxol.

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Example 4: Isolation of mRNA and Northern analysis

Northern and PCR analysis were used to examine the expression level of tubulin mRNAs in *P. microspora* Ne32 and *P. ultimum*.

Four agar plugs of *P. ultimum* or *P. microspora* were inoculated into 40 ml of FM1 medium (5 grams soytone, 5 grams dextrose, 20 grams sucrose, and 1 gram yeast extract per liter of culture) or modified M1D (Li, et al. 1998. *Mycol Res* 102:461-464), respectively, and grown at 24°C without shaking in 250 ml Erlenmeyer flasks. Mycelia were harvested and blotted dry with Whatman paper. One gram (1 g) of mycelia was ground to powder in the presence of liquid nitrogen using mortar and pestle. The powder was transferred to a Dounce homogenizer containing 10 ml of Trizol (GibcoBRL) and homogenized. Total RNA was isolated according to the manufacturer's instructions. Total RNA was dissolved in diethylcarbonate-treated water at room temperature and stored at -80°C. Northern blotting was performed under stringent condition according to standard protocols (Sambrook et al. 1989. *Molecular Cloning: A Laboratory Manual*. Cold Spring Harbor Laboratory, Cold Spring Harbor, NY).

Northern analysis revealed a very low level of beta-tubulin transcripts in *P. microspora* (data not shown). Therefore, total RNA from mycelia grown for 2, 5, 6, or 11 days were converted to cDNA, and used as a template in PCR reactions. Using genespecific primers NETUB5 and NETUB6, a 413 bp beta-tubulin cDNA fragment was amplified from mycelia grown in log (2, 5 and 6 days) or stationary phase (11 days), but not from a control reaction that contained no template.

In *P. ultimum*, total RNA was isolated from mycelia grown for 2, 4, 6, 7, 8 and 10 days, and used in Northern analysis. The beta-tubulin probe hybridized to two transcripts of 1.5 and 1.6 kb, consistent with the sizes of the two cDNAs isolated. The 1.6 kb transcript was present in greater abundance.

Example 5: Binding of [3H]taxol to fungal cells

In animal systems, binding of [³H]taxol to intact cells has been used to characterize interactions between taxol and microtubules (Manfredi, et al. 1982. "Taxol binds to cellular microtubules," *J Cell Biol* 94:688-696). To determine whether the differential taxol sensitivity of *P. microspora* and *P. ultimum* correlates with the taxol binding properties of

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their microtubules, we examined the ability of these fungal cells to specifically bind to [³H]taxol. The total binding of [³H]taxol to fungal cells consists of specific binding to microtubules and nonspecific binding to other cellular structures. The values of total and nonspecific binding were determined by binding of [³H]taxol to fungal cells in the absence or presence of 100-fold excess unlabeled taxol. The specific binding of [³H]taxol was then calculated as the difference between the amount of total and nonspecific binding.

Fresh mycelia from *P. microspora* Ne32, and *P. ultimum* were grown in 140 milliliters modified M1D media in Roux bottles at 24°C for 1-2 days. These actively growing mycelia were transferred to 50 milliliter conical tubes, and centrifuged at 7,000 rpm for 5 minutes at room temperature. Mycelia were suspended in 1 milliliter remaining M1D medium and 1 milliliter fresh M1D medium. Cells were either untreated or pretreated with the anti-mitotic drug thiabendazole to depolymerize microtubules. In pretreated cells, thiabendazole (in DMSO) was added to desired concentrations, and DMSO was adjusted to the same concentration in all samples. Samples were then incubated at room temperature for 3 hours. Subsequently, [³H]taxol (3.7 X 10⁷ Bq/ml, Moravek) was added to desired concentrations either in the presence or absence of 100-fold excess unlabeled taxol. Samples were incubated for 2 hours at room temperature, then quenched on ice. [³H]taxol binding to *P. microspora* cells was performed in the presence of 0.1% (v/v) Triton X-100 to disrupt the cell membrane.

Each GFC filter (Whatman; Clifton, NJ) was weighed using an analytical balance. For *P. ultimum*, mycelia were transferred from conical tubes onto GFC filter hold by a 3-piece filter funnel (Whatman). Conical tubes were washed three times with 10 milliliters of MilliQ H₂O (Millipore, Inc.; Bedford, MA). Mycelia on GFC filter were washed with 120 milliliters of MilliQ H₂O. For *P. microspora*, mycelia were collected by centrifugation at 5,000 rpm for 5 minutes at room temperature, and washed three times with 40 milliliters of MilliQ H₂O. Mycelia were transferred onto GFC filter, and combined with residual mycelia after rinsing conical tubes with 5 milliliters of MilliQ H₂O. GFC filters were dried at 80°C in an oven overnight and then weighed to obtain mycelia dry weight. Filters were counted for 5 minutes under 20 milliliters of Cytoscint (Fisher Scientific; Pittsburgh, PA) in a Beckman LS3801 scintillation counter. Specific binding was calculated as the difference between [³H]taxol bound in the presence and absence of a 100-fold excess unlabeled taxol.

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Nonspecific binding was determined as binding in the presence of 100-fold excess unlabeled taxol.

[³H]taxol was found to bind specifically to *P. ultimum* cells, and the amount of specific binding increased as a function of [³H]taxol concentration (Fig. 7A). In addition, in cells pretreated with thiabendazole to reduce the amount of assembled microtubules, the specific binding of [³H]taxol decreased in a dose-dependent manner (Fig. 7B). In fact, treatment with 1 mM of thiabendazole completely abolished the specific binding of [³H]taxol. These results indicate that taxol is able to interact with *P. ultimum* microtubules, and are consistent with the fact that this organism is sensitive to taxol.

On the other hand, initial experiments showed very low amount of specific binding of [3H]taxol to P. microspora (data not shown). This result could be due to inefficient interactions between taxol and P. microspora microtubules, or alternatively due to a membrane barrier which prevents intracellular accumulation of [3H]taxol. In animal cells, taxol crosses the cell membrane by diffusion due to its hydrophobic character (Manfredi et al. 1982. J Cell Biol 94:688-696). In some cases, resistance to taxol has been associated with P-glycoprotein, a membrane-located pump which causes drug efflux (Jachez et al. 1993. "Restoration of taxol sensitivity of multidrug-resistant cells by the cyclosporine SDZ PSC 833 and the cyclopeptide SDZ 280-446," J Natl Cancer Inst 85:478-483)). There is no information available as to whether P. microspora has such system. It has been shown that treatment of animal cells with nonionic detergents such as 0.1% (v/v) NP-40 or Triton X-100 disrupt the cell membrane, release most soluble proteins including unassembled tubulins, but leave assembled microtubules intact (Schliwa et al. 1981. "Stabilization of the cytoplasmic ground substance in detergent-opened cells and a structural and biochemical analysis of its composition," Proc Natl Acad Sci USA 78:4329-4333; Duerr et al. 1981. "Molecular analysis of cytoplasmic microtubules in situ: identification of both widespread and specific proteins," Cell 24:203-222; Manfredi et al. 1982. J Cell Biol 94:688-696). To evaluate whether a membrane barrier was responsible for the low specific binding, [3H]taxol binding to P. microspora cells in the presence of Triton X-100 was performed. P. microspora cells treated with Triton X-100 (0.1% (v/v)) showed none or very little specific binding of [3H]taxol up to 75 nM [3H]taxol (Fig. 7A). Furthermore, cells pretreated with thiabendazole also showed no specific binding of [3H]taxol in the presence of Triton X-100

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(data not shown). These results indicate that taxol is unable to interact or interacts poorly with microtubules of P. microspora, and are consistent with the fact that this organism is resistant to taxol. In summary, the [3 H]taxol binding results demonstrate that the properties of β -tubulin in these organisms determine their differential sensitivity to taxol.

Taxol stabilizes MTs by binding to beta-tubulin in assembled MTs, and its binding site has been characterized by photo cross-linking, electron crystallography, and mutagenesis. Regions between Amino Acids 1-31 and 217-231 were found to cross-link to the C-3' and C-2 group of taxol, respectively (Rao, et al. 1994. J Biol Chem 269:3132-3134; and Rao, et al. 1995. J Biol Chem 270:20235-20238). Recently, the structure of the betatubulin dimer was solved by electron crystallography of zinc induced sheets of tubulin dimer (Nogales, et al. 1998. Nature 391:199-203). Modeling of taxol bound to this structure shows that the C-3' group of taxol is near Amino Acids 15-25 of beta-tubulin (near the top of helix H1), and the C-2 group is near Amino Acids 212-222 (near helix H6 and the loop between H6-H7). The identification of Amino Acids 15-25 and 217-222 in both cross-linking and electron crystallography studies indicate these regions are important for taxol binding. In addition, the electron crystallography model also shows that Leu273 of bovine beta-tubulin (located in the M-loop) contacts the taxane ring of taxol (Nogales, et al. 1998. Nature 391:199-203). In addition, mutations at Phe270 or Ala364 in the M40 isotype of betatubulin result in taxol resistance in human ovarian cells (Giannakakou, et al. 1997. J Biol Chem 272:17118-17125).

Since the Amino Acids 270, 273 and 364 (marked by # in Fig. 6) do not differ among the fungal beta-tubulins listed in Fig. 6, they are not responsible for the differential taxol response among these organisms. However, comparison of Amino Acids 1-31 and 212-231 (defined here as taxol binding region I and II, respectively) from beta-tubulins of organisms that are taxol-resistant or taxol-sensitive reveal residues that are important for taxol interaction. Fig. 8 provides a comparison of the taxol binding region I and taxol binding region II amino acid sequences for pig (I, SEQ ID NO:34; II, SEQ ID NO:35), human β2 (I, SEQ ID NO:36; II, SEQ ID NO:37), *Drosophila* β1 (I, SEQ ID NO:38; II, SEQ ID NO:39), *Xenopus* β4 (I, SEQ ID NO:40; II, SEQ ID NO:41), *Tetrahymena* (I, SEQ ID NO:42; II, SEQ ID NO:43), *Physarum* β1 (I, SEQ ID NO:44; II, SEQ ID NO:45), *P. ultimum* (I, SEQ ID NO:46; II, SEQ ID NO:47), *P. cinnamomi* (I, SEQ ID NO: 48; II, SEQ

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ID NO: 49) A. klebsiana(I, SEQ ID NO:50; II, SEQ ID NO:51), P. microspora (I, SEQ ID NO:52; II, SEQ ID NO:53), A. nidulans benA (I, SEQ ID NO:54; II, SEQ ID NO:55), and S. cerevisiae (I, SEQ ID NO:56; II, SEQ ID NO:57).

Beta-tubulins from taxol-sensitive organisms such as human, pig, *Drosophila*, *Xenopus, Tetrahymena* and *Physarum* are highly conserved in taxol binding region I and II, and are identical between Amino Acids 15-25 and 217-222 (except a conserved substitution at Amino Acid 23 in *Drosophila* β1). Beta-tubulin from *P. ultimum* displays only four substitutions compared to the above sequences, none of which occurs between Amino Acids 15-25 and 217-222. This similarity is consistent with the fact that *P. ultimum*, like the animal organisms noted above, is taxol-sensitive. Also consistent with this, previous biochemical studies of animal tubulins and data of [³H]taxol binding to *P. ultimum* demonstrated herein (Fig. 7A and 7B), show that taxol binds beta-tubulin in assembled MTs of these organisms (Kellogg, et al. 1989. *J Cell Biol* 109:2977-2991; and Manfredi, J.J. and Horwitz, S.B. 1984. *Pharmacol Ther* 25:83-125). Beta-tubulin sequences from *P. ultimum* and *A. klebsiana* are identical in taxol binding region I and II except Amino Acid 219, but *A. klebsiana* is relatively resistant to taxol (IC₅₀ > 11.7 μM). This reduced sensitivity is due in part to the fact that *A. klebsiana* contains an asparagine at Amino Acid 219, whereas *P. ultimum*, and six other beta-tubulins from taxol-sensitive organisms, have threonine.

Beta-tubulins from taxol-resistant organisms such as *P. microspora*, *A. nidulans* and *S. cerevisiae* are similar to each other within taxol binding region I and II, but differ from the above discussed sequences in seven positions (19, 22, 23, 25, 218, 219, and 221) within regions 15-25 and 217-222. The [³H]taxol binding data presented herein (Fig. 7A and 7B), together with previous biochemical studies (Yoon, Y. and Oakley, B.R. 1995. *Biochem* 34:6373-6381; and Bames, et al. 1992. *Mol Biol Cell* 3:29-47), show that beta-tubulins in assembled MTs of these organisms are unable to efficiently bind taxol. These sequences contain the asparagine (or glutamine in the case of *S. cerevisiae*) at Amino Acid 219, as observed in *A. klebsiana*, a substitution that contributes in part to the reduced sensitivity to taxol in these fungi. Other substitutions, including which involve differences in charge and polarity such as changes from Lys19 to Ala, Glu22 to Gln, and Val23 to Thr, also contribute to the taxol resistant phenotype of these organisms.

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These results indicate that a number of residues including threonine at Amino Acid 219 are important in the binding of beta-tubulin to taxol. In particular, Amino Acid 219 plays an important role in determining taxol binding property of beta-tubulin and, consequently, the taxol-sensitivity of cells. Beta-tubulins from taxol-sensitive species have Thr219 (threonine at Amino Acid 219), while those from taxol-resistant species have Asn219 (asparagine at Amino Acid 219) or Glu219 (glutamine at Amino Acid 219). The taxol sensitivity of *P. cinnamomi* is consistent with the presence of Thr219 in TUBB-pc (SEQ ID NO:6) and not Asn219 as previously reported by Weerakoon et al. The presence of Asn219 (asparagine at Amino Acid 219) found in *P. microspora* is consistent with the taxol resistance of this species. Using the information that the presence of threonine at Amino Acid 219 in beta-tubulins corresponds to taxol-binding and taxol-sensitivity, taxol analogs or other compounds can be designed which mimic the interaction of taxol with beta-tubulin. Further, such information can also be used to generate mutant beta-tubulins resistant to taxol by substituting the threonine for another amino acid residue at Amino Acid 219.

Example 6: Sensitivity to microtubule-depolymerization drugs.

The effect of several MT-depolymerization drugs on the growth of *P. microspora* Ne32, *P. ultimum* and *A. klebsiana* was examined. These drugs included colchicine, colcemid (a synthetic derivative of colchicine), and two benzimidazole drugs, nocodazole and thiabendazole.

Colchicine, colcemid, nocodazole, and thiabendazole were obtained from Sigma Chemical Company (St. Louis, MO). A stock solution of colchicine was prepared in water, and other stock solutions in DMSO. An agar plug (6 mm in diameter) of fresh mycelia was transferred onto PDA plates containing 1 % (v/v) DMSO in the presence or absence of an anti-microtubule agent. Fungal colonies were grown at 24°C for 24 hours in the case of *P. ultimum* or 48 hours in the case of *P. microspora* and *A. klebsiana*. The growth inhibitory effect of these anti-mitotic agents was measured by the size of colony diameters.

Biochemical and genetic evidence has shown that these drugs bind to beta-tubulin in the tubulin dimer and cause MT depolymerization (Davidse, L.C. and Flach, W. 1978. "Interaction of thiabendazole with fungal tubulin," *Biochim Biophys Acta* 543:82-90; Jung,

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M.K. and Oakley, B.R. 1990. "Identification of an amino acid substitution in the β-tubulin gene of Aspergillus nidulans that confers thiabendazole resistance and benomyl supersensitivity," Cell Motil Cytoskeleton 11:87-94; Manfredi, J.J. and Horwitz, S.B. 1984. Pharmacol Ther 25:83-125). It has been shown that many fungi are resistant to colchicine (Cameron et al. 1990. J Biol Chem 265:15245-15252; Kilmartin, J.V. 1981. "Purification of yeast tubulin by self-assembly in vitro," Biochem 20:3629-3633; and Davidse, L.C. and Flach, W. 1977. "Differential binding of methyl benzimidazole-2-yl-carbarnate to fungal tubulin as a mechanism of resistance to this antimitotic agent in mutant strains of Aspergillus nidulans," J Cell Biol 72:174-193), but are sensitive to nocodazole (Kilmartin, J.V. 1981. Biochem 20:3629-3633) and thiabendazole (Davidse, L.C. and Flach, W. 1978. Biochim Biophys Acta 543:82-90).

As shown in Table II, the three fungal species tested herein were resistant to colchicine and colcemid (IC₅₀ >100 μ M), and were sensitive to nocodazole (IC₅₀ 2-22 μ M). These results are consistent with the studies noted above. In contrast, these fungi were differentially sensitive to thiabendazole. *P. microspora* was highly sensitive (IC₅₀ 3 μ M), while *P. ultimum* and *A. klebsiana* were less sensitive (IC₅₀ 270-350 μ M). These results demonstrate that the biochemical properties of beta-tubulin differ in these three fungi.

Table II: Sensitivity of fungi to microtubule depolymerization drugs

Drug	P. microspora	P. ultimum	A. klebsiana
colchicine	>100	>100	>100
colcemid	>100	>100	>100
nocodazole	2	22	2
thiabendazole	3	350	270

Mutations at Amino Acids 6, 165, 167, 198, 200 and 241 in beta-tubulin (marked by asterisks in Fig. 6) result in altered sensitivity to thiabendazole and other benzimidazole drugs in yeast, *N. crassa*, *A. nidulans* benA, and *Trichoderma viride* (Thomas, et al. 1985. "Isolation and characterization of mutations in the β-tubulin gene of *Saccharomyces* cerevisiae," Genetics 112:715-734; Orbach, et al. 1986. Mol Cell Biol 6:2452-2461; Jung, et

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al. 1992. "Amino acid alterations in the β-tubulin gene of Aspergillus nidulans that confer benomyl resistance," Cell Motil Cytoskeleton 22:170-174; Jung, M.K. and Oakley, B.R. 1990. Cell Motil Cytoskeleton 17:87-94; Fugimura, et al. 1992. "A single amino-acid substitution in the beta-tubulin gene of Neurospora confers both cabendazim resistance and diethofencarb sensitivity," Curr Genet 21:399-404; and Goldman et al. 1993. Mol Gen Genet 240:73-80). These six residues in beta-tubulin from P. microspora are identical to those observed in beta-tubulin from other thiabendazole sensitive species such as N. crassa and A. nidulans benA. In contrast, beta-tubulin from P. ultimum and A. klebsiana differ at Amino Acids 165, 167 and 200. It has been previously shown that a phenylalanine-to-tyrosine change at Amino Acid 167 results in benzimidazole resistance in N. crassa (Orbach, et al. 1986. Mol Cell Biol 6:2452-2461), and the fact that both P. ultimum and A. klebsiana have a tyrosine at this position accounts for their resistance to such drugs. In summary, the differential sensitivity to thiabendazole exhibited by these three fungi is consistent with the comparison of fungal beta-tubulins shown in Fig. 6.

Example 7: Production of Antibodies Capable of Distinguishing Taxol-Binding and Non-Binding Beta-tubulins

Monoclonal or polyclonal antibodies can be raised against the following antigens: 1) native beta-tubulins extracted from *P. microspora*, *P. ultimum*, or *P. cinnamomi*; 2) beta-tubulins of *P. microspora*, *P. ultimum*, or *P. cinnamomi* produced from a heterologous system such as *E. coli*, yeast, and insect cells; and 3) synthetic peptide corresponding to the SEQ ID NO:2, SEQ ID NO:4 or SEQ ID NO:6, and preferably comprising at least one taxol binding region. The antibodies are used to interact with the above mentioned beta-tubulins using Elisa or Western blotting using standard protocols (Harlow, E.D. and Lane, D. 1988. *Antibodies: A Laboratory Manual*). The antibodies which could distinguish the taxol binding beta-tubulin from the taxol non-binding beta-tubulin are selected as the reagent.

A specific example is to raise polyclonal or monoclonal antibodies to synthetic peptides corresponding to SEQ ID NO:4 or SEQ ID NO:6 which comprise at least one taxol binding region, for instance containing the taxol-binding region II comprising Thr219 or in which the Thr219 is replaced by Asn219/Gln219. The ability of these antibodies to interact with beta-tubulin is examined using Elisa using standard protocols. The antibody which can binds to peptide containing Thr219 but not to peptide containing Asn/Gln 219 is selected as

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the reagent which is specific for the taxol-binding site containing Thr 219. On the other hand, the antibody which specifically binds to the peptide containing Asn219/Gln 219 but not to the peptide containing Thr 219 is selected as the reagent which specifically recognizes taxol binding site devoid of Thr 219.

Example 8: Screening Assays to Detect Beta-Tubulin in Matter

Several assays can be used to determine if a composition of matter contains betatubulin capable of binding taxol. These assays are useful for screening a variety of compositions of matter, including living matter such as plants or microorganisms, or nonliving matter such as plant materials or patient samples for the presence of beta-tubulin.

The first assay is performed using Northern or Southern hybridization method well known in the art (Sambrook, et al. *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989). The total RNA, mRNA or genomic DNA are isolated from the composition of matter and separated by electrophoresis. DNA, synthetic oligonucleotide, or RNA corresponding to the coding region or a portion of beta-tubulin (e.g., derived from SEQ ID NO:1, SEQ ID NO: 3 or SEQ ID NO:5) which comprises at least one taxol binding region will be used to synthesize isotopically labeled probes. Hybridization with a probe derived from SEQ ID NO:1 will indicate beta-tubulin with high probability of taxol resistance. On the other hand, the hybridization with a probe derived from SEQ ID NO:3 or SEQ ID NO:5 will indicate beta-tubulin with a high probability of taxol sensitivity.

The second assay is to use a PCR-based assay using standard protocols (Sambrook, et al. *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989). Both genomic DNA or cDNA converted from total RNA or mRNA are used as template in a PCR assay. Gene-specific or degenerate primers corresponding to the coding region of beta-tubulin (e.g., derived from SEQ ID NO:1, SEQ ID NO:3 or SEQ ID NO:5) which comprises at least one taxol binding region will be synthesized. Only DNA containing the appropriate primer sequences will be amplified, and all other variations will be suppressed. The amplification of PCR fragment of the predicted size using primers derived from SEQ ID NO:3 or SEQ ID NO:5 but not from primers derived from SEQ ID NO:1 will indicate high probability of taxol binding beta-tubulin. On

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the other hand, the amplification of a PCR fragment of the predicted size using primers derived from SEQ ID NO:1 but not from primers derived from SEQ ID NO:3 or SEQ ID NO:5 will indicate high probability of taxol non-binding beta-tubulin. The subsequent obtaining of the beta-tubulin sequence and examination of the presence or absence of Thr219 residue will provide further determination.

The third assay is to use Elisa or Western blotting using standard protocols (Harlow, E.D. and Lane, D. 1988. *Antibodies: A Laboratory Manual*). Cell extracts of the composition of matter are prepared. Synthetic peptide, or native beta-tubulins extracted from *P. microspora*, *P. ultimum*, or *P. cinnamomi*, or produced from a heterologous system such as *E. coli*, yeast, and insect cells will be used to raise polyclonal or monoclonal antibodies. The antibodies will be used in the above mentioned Elisa or Western blotting. The antibody which recognizes the taxol binding from the non taxol binding is used in these assays.

Example 9: Construction of Taxol-sensitive and Taxol-resistant Isogenic Strains

P. ultimum contains a single beta-tubulin. In vitro, its beta-tubulin gene or cDNA can be altered to change the Thr219 to a different residue, for instance to Asn219 or Gln219. This altered DNA sequence is cloned into a transformation vector, and used to transform the wild-type strain P. ultimum using established protocols (Balance, et al. 1985. Gene 36:321-331). Homologous recombination between the wild-type beta-tubulin gene and the modified beta-tubulin in the vector occur. Transformed fungus are selected on media containing taxol. The taxol-resistant clones are selected and their beta-tubulin cDNA sequenced to confirm the absence of Thr 219. The taxol-resistant isogenic strain of P. cinnamomi is similarly constructed and used in screening assays as described in later examples. The only difference between these isogenic strains is that the taxol-sensitive strain is capable of binding to taxol due to the presence of Thr 219, and the taxol-resistant strain is incapable of binding to taxol due to the absence of Thr 219. Such taxol-resistant strains can be used in combination with the wild-type taxol-sensitive strains for screening as described in later examples.

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Several assays can be used to detect taxol or taxol-like compounds in a composition of matter. These assays are useful for screening a variety of compositions of matter, including living matter such as plants or microorganisms, or non-living matter such as plant materials, patient samples, or compound libraries for the presence of taxol or taxol-like compounds.

One screening method is to use taxol-resistant *P. microspora* in combination with the taxol-sensitive *P. ultimum* or *P. cinnamomi*. Taxol inhibits the growth of both *P. ultimum* by binding to their beta-tubulin, while taxol does not affect the growth of *P. microspora* since it does not interact with its beta-tubulin. A composition of matter which is capable of the inhibition of *P. ultimum*, but not *P. microspora* has a high probability of containing taxol-or a taxol-like compound.

An improved screening method uses taxol-sensitive and taxol-resistant isogenic strains of *P. ultimum* or *P. cinnamomi* as described in above example. The composition of matter is used to examine its effect on the growth of both the taxol-sensitive as well as the taxol-resistant strains. The inhibition of the taxol-sensitive strain but not the taxol-resistant strain indicates the presence of taxol or a taxol-like compound. On the other hand, the non-inhibition of both the taxol-sensitive and taxol-resistant strains indicates the absence of taxol or a taxol-like compound.

Composition of matter can be screened for the presence of taxol or taxol-like compounds based on their ability to promote the assembly of microtubules, as well as to stabilize assembled microtubules in conditions such as cold which otherwise cause depolymerization (Schiff, et al. 1979; Horwitz, 1981). The alpha- and beta-tubulins used in these assays can be from the following sources. 1) native microtubules consisting of beta-tubulins and alpha-tubulins extracted from *P. ultimum* or *P. cinnamomi*; 2) beta-tubulins extracted from *P. ultimum* or *P. cinnamomi*; 3) all or portions of SEQ ID NO:4 or SEQ ID NO:6 produced from a heterologous system such as *E. coli*, yeast, insect cells or the like and alpha-tubulin either from *P. ultimum*, *P. cinnamomi* or another source. If the composition matter has the ability to promote the assembly of these MTs, as well as to prevent

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depolymerization of assembled MTs in conditions which otherwise cause depolymerization, the composition of matter is likely to contain taxol or a taxol-like compound. Meanwhile, these isolated compounds should be unable to promote the assembly of MTs as well as prevent the depolymerization of MTs which consist of beta-tubulin derived from *P*. *microspora*.

An alternative screening method can be performed based on the competitive inhibition of [³H]taxol binding to MTs in P. ultimum or P. cinnamomi by taxol or taxol-like compounds. The specific binding of [³H]taxol to P. ultimum is performed as described in Example 5. The amount of [³H]taxol specifically bound to P. ultimum in the absence of inhibitors is considered 100%. The composition of matter is added to the assay mixture, and the amount of [³H]taxol specifically bound to P. ultimum in the presence of the composition of matter is measured. Reduction in the [³H]taxol specific binding indicates that the composition of matter possesses taxol-like quality. If increased concentrations of the composition of matter can completely inhibit the [³H]taxol binding, it will indicate that the compound likely binds to the same binding site in the beta-tubulin in MTs.

The screening of compositions of matter for taxol or taxol-like compounds can be performed by one of the above methods. Preferably, one of the first two methods is used for an initial screening, since they are simple to perform and easily handle large amounts of samples. The third and fourth method can be used for subsequent screening.

Example 11: Screening Assay to Distinguish Taxol-Sensitivity of A Patient Sample

In this diagonostic assay, antibodies depicted in Example 7 which could distinguish taxol-binding beta-tubulin from the non-binding beta-tubulin are used. Cellular proteins are extracted from a tumor specimen from a patient sample to detect the presence of a beta-tubulin with either taxol-binding or non-binding capabilities.

For example, in a diagnostic assay to screen for tumors resistant to taxol, the taxol binding regions of taxol-sensitive and taxol-resistant beta-tubulins of the present invention (e.g., SEQ ID NO:2, SEQ ID NO:4, and SEQ ID NO:6) are used to raise monoclonal or polyclonal antibodies using standard methods well known in the art (Harlow, E.D. and Lane, D. 1988. *Antibodies: A Laboratory Manual*). For example, monoclonal antibody probes are

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reacted with a patient sample, such as a tumor specimen, to detect the presence of a beta-tubulins with either taxol-binding or non-binding capabilities. Visualization of antibody-antigen binding is mediated through any means known in the art, e.g., secondary radiolabeled or fluorescent antibodies or colorimetric methods using peroxidase and/or alkaline phosphatase (Harlow, E.D. and Lane, D. 1988. *Antibodies: A Laboratory Manual*). The detection of beta-tubulins with taxol-binding capability, i.e., taxol-sensitive beta-tubulins, corresponds to a positive response to taxol therapy. Alternatively, the detection of non-binding taxol-resistant beta-tubulins and/or the absence of taxol-sensitive beta-tubulins corresponds to a diminished or lack of response to taxol therapy.

Example 12: Biocontrol of Taxol-sensitive Pathogenic Oomycetes Using *P. microspora* on Plants

Many oomycetes including *P. ultimum* and *P. cinnamomi* are plant pathogens which can cause crop damage and result in severe economical loss. For instance, *P. ultimum* causes root rot of beans, and *P. cinnamomi* causes root rot of Avacado (ATCC: Catalogue of Filamentous Fungi, 18th edition, 1991). Many of the oomycetes are also taxol-sensitive (Young, et al. 1992. "Antifungal properties of taxol and various analogues," *Experientia* 48:882-885). Two of these strains, *P. ultimum* and *P. cinnamomi*, contain threonine at Amino Acid 219.

The biocontrol method of the present invention involves a two-step process: 1) the taxol sensitivity of the plant pathogen is determined and 2) if the plant pathogen is taxol-sensitive, a taxol-producing *P. microspora* is applied to the infected plants and surrounding soil as a source of growth-inhibiting taxol.

The taxol sensitivity of the plant pathogen is first determined. One method of identifying taxol sensitivity is to determine the presence or absence of threonine at Amino Acid 219. If the identity of the pathogen is known, DNA and protein databases are searched to determine whether the beta-tubulin sequence has been reported, if so, the identity of Amino Acid 219 is determined from the database. If the pathogen's beta-tubulin sequence is unavailable, the cDNA sequence is isolated and analyzed to determine the identity of Amino Acid 219. The presence of threonine at Amino Acid 219 in the pathogen's beta-tubulin gene indicates sensitivity to taxol, and thus, the pathogen is designated as treatable by a taxol-

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producing *P. microspora*. If Amino Acid 219 is not threonine, the taxol sensitivity would have to be determined by other means such as taxol growth inhibition. Other screening methods presented herein for determining the presence of taxol-binding beta-tubulins can also be used.

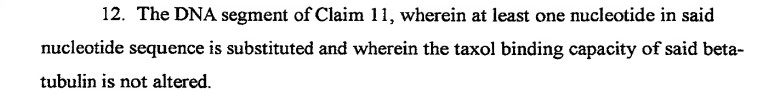
It has been previously reported that *P. microspora* produces taxol at 50 ug /liter (Strobel, et al. 1996. *Microbiol* 142:435-440) and secrets taxol outside of the fungal cell. At this taxol concentration, the growth of *P. ultimum* and *P. cinnamomi* is inhibited (see Fig. 1). For treatment, *P. microspora* is inoculated to the area of plants or soil infected with the taxol-sensitive pathogen. The growth of *P. microspora* results in the secretion of taxol, which consequently inhibits the growth of these taxol-sensitive organisms.

Example 13: Use of Crystal Structures in Design of Antineoplastic or Antifungal Drugs

The three-dimensional structure of beta-tubulins are used to rationally design taxol-like compounds using methods known in the art (Ealick, et al. 1991. "Application of crystallographic and modeling methods in the design of purine nucleoside phosphorylase inhibitors," *Science* 88:11540-11544; Rossman, et al. 1991. "Application of crystallography to the design of antiviral agents," *Infectious Agents and Disease* 1:3-10). As taught by the present invention, application of the knowledge that Thr219 in the protein structure plays an important role in binding of taxol to taxol-like compounds can be critically applied to the development of drugs having taxol-like activities.

We claim:

- 1. A purified DNA segment encoding a beta-tubulin of the fungal species Pestalotiopsis microspora or a portion thereof.
- 2. The DNA segment of Claim 1, wherein said portion encodes at least one taxol binding site.
- 3. The DNA segment of Claim 2, wherein said portion encodes a protein having taxol binding site I and taxol binding site II.
- 4. The DNA segment of Claim 3, wherein said protein is able to interact with alphatubulin.
- 5. The DNA segment of Claim 1, wherein said DNA segment comprises at least a portion of SEQ ID NO:1.
- 6. The DNA segment of Claim 5, wherein said portion comprises the nucleotide sequence from nucleotide 75 through nucleotide 167 of SEQ ID NO:1.
- 7. The DNA segment of Claim 6, wherein at least one nucleotide in said nucleotide sequence is substituted.
- 8. The DNA segment of Claim 5, wherein said portion comprises the nucleotide sequence from nucleotide 708 through nucleotide 764 of SEQ ID NO:1
- 9. The DNA segment of Claim 8, wherein at least one nucleotide in said nucleotide sequence is substituted.
- 10. The DNA segment of Claim 9, wherein nucleotide 729, nucleotide 730, nucleotide 731 or mixtures thereof are substituted.
- 11. The DNA segment of Claim 5, comprising the nucleotide sequence from nucleotide 75 to nucleotide 1412 of SEQ ID NO:1, said DNA segment encoding a betatubulin.



- 13. The DNA segment of Claim 11, wherein at least one nucleotide in said nucleotide sequence is substituted and wherein the taxol binding capacity of said betatubulin is altered.
- 14. An amino acid sequence comprising at least a portion of a beta-tubulin of the fungal species *Pestalotiopsis microspora*.
- 15. The amino acid sequence of Claim 14, wherein said portion comprises at least one taxol binding site.
- 16. The amino acid sequence of Claim 15, wherein said portion comprises taxol binding site I and taxol binding site II.
- 17. The amino acid sequence of Claim 16, wherein said portion is able to interact with alpha-tubulin.
- 18. The amino acid sequence of Claim 14, wherein said amino acid sequence comprises at least a portion of the beta-tubulin as depicted in SEQ ID NO:2.
- 19. The amino acid sequence of Claim 18, wherein said portion comprises Amino Acids 1-31 of SEQ ID NO:2.
- 20. The amino acid sequence of Claim 19 having at least one amino acid substitution.
- 21. The amino acid sequence of Claim 18, wherein said portion comprises Amino Acids 212-230 of SEQ ID NO:2.
- 22. The amino acid sequence of Claim 21 having at least one amino acid substitution.
- 23. The amino acid sequence of Claim 18, wherein said portion comprises an amino acid substitution at Amino Acid 219.

- 24. The amino acid sequence of Claim 18, wherein said portion consists essentially of Amino Acids 1-446 of SEQ ID NO:2 and wherein said portion behaves as a taxol-resistant beta-tubulin.
- 25. The amino acid sequence of Claim 24, wherein said portion contains at least one amino acid substitution that alters the taxol binding property of said portion.
- 26. The amino acid sequence of Claim 24, wherein said portion contains at least one amino acid substitution that does not alter the taxol binding property of said portion.
- 27. The amino acid sequence of Claim 14, wherein said amino acid sequence is substituted with any amino acid which perturbs the three-dimensional structure of said amino acid sequence surrounding Amino Acid 219 as numbered in SEQ ID NO:2.
- 28. A purified DNA segment encoding a beta-tubulin of the fungal species *Pythium* ultimum or a portion thereof.
- 29. The DNA segment of Claim 28, wherein said portion encodes at least one taxol binding site.
- 30. The DNA segment of Claim 29, wherein said portion encodes a protein having taxol binding site I and taxol binding site II.
- 31. The DNA segment of Claim 30, wherein said protein is able to interact with alpha-tubulin.
- 32. The DNA segment of Claim 28, wherein said DNA segment comprises at least a portion of SEQ ID NO:3.
- 33. The DNA segment of Claim 32, wherein said portion comprises the nucleotide sequence from nucleotide 92 through nucleotide 184 of SEQ ID NO:3.
- 34. The DNA segment of Claim 33, wherein at least one nucleotide in said nucleotide sequence is substituted.
- 35. The DNA segment of Claim 32, wherein said portion comprises the nucleotide sequence from nucleotide 725 through nucleotide 781 of SEQ ID NO:3

- 36. The DNA segment of Claim 35, wherein at least one nucleotide in said nucleotide sequence is substituted.
- 37. The DNA segment of Claim 35, wherein nucleotide 746, nucleotide 747, nucleotide 748 or mixtures thereof are substituted.
- 38. The DNA segment of Claim 32, comprising the nucleotide sequence from nucleotide 92 to nucleotide 1429 of SEQ ID NO:3, said DNA segment encoding a betatubulin.
- 39. The DNA segment of Claim 38, wherein at least one nucleotide in said nucleotide sequence is substituted and wherein the taxol binding capacity of said betatubulin is not altered.
- 40. The DNA segment of Claim 38, wherein at least one nucleotide in said nucleotide sequence is substituted and wherein the taxol binding capacity of said betatubulin is altered.
- 41. An amino acid sequence comprising at least a portion of a beta-tubulin of the fungal species *Pythium ultimum*.
- 42. The amino acid sequence of Claim 41, wherein said amino acid sequence comprises at least one taxol binding site.
- 43. The amino acid sequence of Claim 42, wherein said portion comprises taxol binding site I and taxol binding site II.
- 44. The amino acid sequence of Claim 43, wherein said portion is able to interact with alpha-tubulin.
- 45. The amino acid sequence of Claim 41, wherein said amino acid sequence comprises at least a portion of the beta-tubulin as depicted in SEQ ID NO:4.
- 46. The amino acid sequence of Claim 45, wherein said portion comprises Amino Acids 1-31 of SEQ ID NO:4

- 47. The amino acid sequence of Claim 46, having at least one amino acid substitution.
- 48. The amino acid sequence of Claim 45, wherein said portion comprises Amino Acids 212-230 of SEQ ID NO:4
- 49. The amino acid sequence of Claim 48, having at least one amino acid substitution.
- 50. The amino acid sequence of Claim 45, wherein said portion comprises an amino acid substitution at Amino Acid 219.
- 51. The amino acid sequence of Claim 45, wherein said portion consists essentially of Amino Acids 1-446 of SEQ ID NO:4 and wherein said portion behaves as a taxol-sensitive beta-tubulin.
- 52. The amino acid sequence of Claim 51, wherein said portion contains at least one amino acid substitution that alters the taxol binding property of said portion.
- 53. The amino acid sequence of Claim 51, wherein said portion contains at least one amino acid substitution that does not alter the taxol binding property of said portion.
- 54. The amino acid sequence of Claim 41, wherein said amino acid sequence is substituted with any amino acid which perturbs the three-dimensional structure of said amino acid sequence surrounding Amino Acid 219 as numbered in SEQ ID NO:4.
- 55. A purified DNA segment encoding a beta-tubulin of the fungal species *Phytophthora cinnamomi* or a portion thereof, wherein said DNA segment consists essentially of at least a portion of SEQ ID NO:5.
- 56. The DNA segment of Claim 55, wherein said portion comprises the nucleotide sequence from nucleotide 11 through nucleotide 103 of SEQ ID NO:5.
- 57. The DNA segment of Claim 56, wherein at least one nucleotide in said nucleotide sequence is substituted, providing that when nucleotide substitution changes only

one amino acid code nucleotide 80 cannot consist of adenine while nucleotide 81 is thymine and nucleotide 82 is adenine, cytosine or thymine.

- 58. The DNA segment of Claim 55, wherein said portion comprises the nucleotide sequence from nucleotide 644 through nucleotide 700 of SEQ ID NO:5
- 59. The DNA segment of Claim 58, wherein at least one nucleotide in said nucleotide sequence is substituted, providing that when nucleotide substitution changes only one amino acid code nucleotide 665 cannot be adenine while nucleotide 666 is adenine and nucleotide 667 is cytosine or thymine.
- 60. The DNA segment of Claim 55, comprising the nucleotide sequence from nucleotide 11 to nucleotide 1342 of SEQ ID NO:5, said DNA segment encoding a betatubulin.
- 61. The DNA segment of Claim 60, wherein at least one nucleotide in said nucleotide sequence is substituted, providing that when nucleotide substitution changes only one amino acid code nucleotide 665 cannot be adenine while nucleotide 666 is adenine and nucleotide 667 is cytosine or thymine.
- 62. The DNA segment of Claim 60 or 61, wherein at least one nucleotide in said nucleotide sequence is substituted, and wherein the taxol binding capacity of said betatubulin is not altered.
- 63. The DNA segment of Claim 60 or 61, wherein at least one nucleotide in said nucleotide sequence is substituted and wherein the taxol binding capacity of said betatubulin is altered.
- 64. An amino acid sequence comprising at least a portion of a beta-tubulin of the fungal species *Phytophthora cinnamomi* as depicted in SEQ ID NO:6.
- 65. The amino acid sequence of Claim 64, wherein said portion comprises Amino Acids 1-31 of SEQ ID NO:6.

- 66. The amino acid sequence of Claim 65, having at least one amino acid is substituted, providing that when only one amino acid is substituted Amino Acid 24 is not isoleucine.
- 67. The amino acid sequence of Claim 64, wherein said portion comprises Amino Acids 212-230 of SEQ ID NO:6.
- 68. The amino acid sequence of Claim 67, having at least one amino acid is substituted, providing that when only one amino acid is substituted Amino Acid 219 is not asparagine.
- 69. The amino acid sequence of Claim 64, wherein said portion comprises an amino acid substitution at Amino Acid 219, wherein said Amino Acid 219 is not substituted with asparagine.
- 70. The amino acid sequence of Claim 64, wherein said portion consists essentially of Amino Acids 1-446 of SEQ ID NO:6 and wherein said portion behaves as a taxol-sensitive beta-tubulin.
- 71. The amino acid sequence of Claim 70, wherein said portion contains at least one amino acid substitution, providing that when only one amino acid is substituted Amino Acid 219 is not asparagine, and wherein said amino acid substitution that alters the taxol binding property of said portion.
- 72. The amino acid sequence of Claim 70, wherein said portion contains at least one amino acid substitution, providing that when only one amino acid is substituted Amino Acid 219 is not asparagine, and wherein said amino acid substitution does not alter the taxol binding property of said portion.
- 73. The amino acid sequence of Claim 64, wherein said amino acid sequence is substituted with any amino acid which perturbs the three-dimensional structure of said amino acid sequence surrounding Amino Acid 219 and wherein when only one amino acid is substituted at Amino Acid 219 said substituted amino acid is not asparagine.
- 74. A vector comprising a purified DNA segment encoding a beta-tubulin of the fungal species *Pestalotiopsis microspora* or a portion thereof.

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- 75. The vector of Claim 74, wherein said portion encodes at least one taxol binding site.
- 76. A vector comprising a purified DNA segment encoding a beta-tubulin of the fungal species *Pythium ultimum* or a portion thereof.
- 77. The vector of Claim 76, wherein said portion encodes at least one taxol binding site.
- 78. A vector comprising a purified DNA segment encoding a beta-tubulin of the fungal species *Phytophthora cinnamomi* wherein said DNA segment consists essentially of SEQ ID No:5 or a portion thereof.
- 79. The vector of Claim 78, wherein said portion encodes at least one taxol binding site.
- 80. A method of determining the taxol binding capacity of a beta-tubulin or betatubulin-like compound comprising

providing antibodies raised against amino acid sequences comprising at least one taxol binding site of a beta-tubulin from a taxol-resistant *Pestalotiopsis microspora*, a taxol-sensitive *Pythium ultimum*, or taxol-sensitive *Phytophthora cinnamomi* as depicted in SEQ ID NO:6 to form a reagent, wherein said antibodies distinguish between taxol-binding and non-taxol-binding properties;

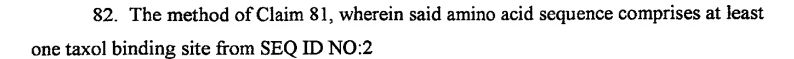
contacting said beta-tubulin with said reagent; and

determining degree of binding between said antibodies in said reagent and said betatubulin or beta-tubulin-like compound;

whereby binding of antibodies which specifically recognize taxol-binding properties indicate taxol sensitive; whereby binding of antibodies which specifically recognize taxol-non-binding properties indicate taxol resistance.

81. The method of Claim 80, wherein said antibodies are raised against an amino acid sequence comprising at least one taxol binding site of a beta-tubulin from a taxol-resistant *Pestalotiopsis microspora*.

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- 83. The method of Claim 80, wherein said antibodies are raised against an amino acid sequence comprising at least one taxol binding site of a beta-tubulin from a taxol-sensitive *Pythium ultimum*.
- 84. The method of Claim 81, wherein said amino acid sequence comprises at least one taxol binding site from SEQ ID NO:4
- 85. The method of Claim 80, wherein said antibodies are raised against an amino acid sequence comprising at least one taxol binding site of a beta-tubulin from a taxol-sensitive *Phytophthora cinnamomi* as depicted in SEQ ID NO:6.
- 86. The method of Claims 80, 81, 82, 83, 84 or 85, wherein said beta-tubulin or beta-tubulin-like protein is selected from the group consisting of recombinantly expressed protein, exogenously isolated protein, synthetic peptides, and cell cultures.
- 87. A method of screening a composition of matter for the presence of taxol or taxol-like compounds comprising

providing beta-tubulins with amino acid sequences comprising both taxol binding sites from taxol-sensitive *Pythium ultimum* or taxol-sensitive *Phytophthora cinnamomi* as depicted in SEQ ID NO:6 in addition to alpha-tubulin from any taxol-sensitive organism to form a reagent;

contacting said composition of matter with said reagent; and

determining the ability of the composition of matter to promote MT assembly or ability to prevent depolymerization of assembled MTs under depolymerizing conditions;

whereby the ability to promote microtubule assembly or prevent depolymerization indicate the possible presence of taxol or taxol-like compounds in said composition of matter.

88. A method of screening a composition of matter for the presence of taxol or taxol-like compounds comprising

providing mycelia of taxol-sensitive *Pythium ultimum* or a taxol-sensitive *Phytophthora cinnamomi* as depicted in SEQ ID NO:6;

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contacting said composition of matter with said mycelia in the presence of said labeled taxol; and

determining degree of competitive inhibition of binding between said beta-tubulins and said labeled taxol by said composition of matter;

whereby the composition of matter is determined to possess taxol or taxol-like compounds if it is able to block taxol binding to the beta-tubulins from the taxol-sensitive *Pythium ultimum* or *Phytophthora cinnamomi*.

89. A method of altering the taxol binding property of a recombinantly expressed beta-tubulin or a portion thereof comprising

determining the identity of the codon at Amino Acid 219 as numbered in SEQ ID NO:2 in the coding region of the vector; and

if said codon at Amino Acid 219 codes for any amino acid except threonine, substituting nucleotides in said codon to code for threonine at Amino Acid 219 to alter a non-taxol-binding beta-tubulin or portion thereof to a taxol-binding beta-tubulin or portion thereof or if said codon at Amino Acid 219 codes for threonine, substituting nucleotides in said codon to code for any amino acid except threonine at Amino Acid 219 to alter a taxol-binding beta-tubulin or portion thereof.

90. A method of developing a taxol-sensitive fungal cell from a taxol-resistant fungal cell comprising

transforming said non-taxol-sensitive fungal cell by introducing a DNA segment encoding taxol-binding beta-tubulin comprising threonine at Amino Acid 219 as numbered in SEQ ID NO:2;

wherein said transformed fungal cell expresses said DNA segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.

91. A transgenic taxol-sensitive fungal cell transformed by introducing a DNA segment encoding taxol-binding beta-tubulin comprising threonine at Amino Acid 219 as numbered in SEQ ID NO:2, wherein said transformed fungal cell expresses said DNA

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segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.

92. A method of developing a taxol-resistant fungal cell from a taxol-sensitive fungal cell comprising

transforming said taxol-sensitive fungal cell by introducing a DNA segment encoding non-taxol-binding beta-tubulin wherein the amino acid at Amino Acid 219 as numbered in SEQ ID NO:2 is not threonine;

wherein said transformed fungal cell over-expresses said DNA segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.

- 93. A transgenic taxol-sensitive fungal cell transformed by introducing a DNA segment encoding taxol-binding beta-tubulin comprising threonine at Amino Acid 219 as numbered in SEQ ID NO:2, wherein said transformed fungal cell over-expresses said DNA segment under the control of a suitable constitutive or inducible promoter when exposed to conditions which permit expression.
- 94. A method of screening a composition of matter for the presence of taxol or taxol-like compounds comprising

providing distinguishable taxol-resistant and taxol-sensitive fungal cells; contacting said composition of matter with said fungal cells; determining the growth inhibition of said fungal cells;

whereby the composition of matter is determined to possess taxol or taxol-like compounds if it is able to inhibit the growth of taxol-sensitive fungal cells but not able to inhibit the growth of taxol-resistant fungal cells.

- 95. The method of Claim 94, wherein said distinguishable taxol-resistant and taxol-sensitive fungal cells consists essentially of transgenic taxol-resistant and taxol-sensitive isogenic fungal cells.
- 96. The method of Claim 94, wherein said taxol-resistant fungal cells are derived from a fungi which is unrelated to the fungi from which the taxol-sensitive fungal cells are derived.

- 97. A method for controlling the growth of plant pathogens comprising determining the taxol sensitivity of said plant pathogen; and if said pathogen is determined to be taxol-sensitive, said plant and soil surrounding said plant are treated with a taxol-producing *P. microspora*.
- 98. The method of Claim 97, wherein the taxol sensitivity of said plant pathogen is determined by identifying Amino Acid 219, wherein the plant is designated as taxol-sensitive if Amino Acid 219 is threonine.

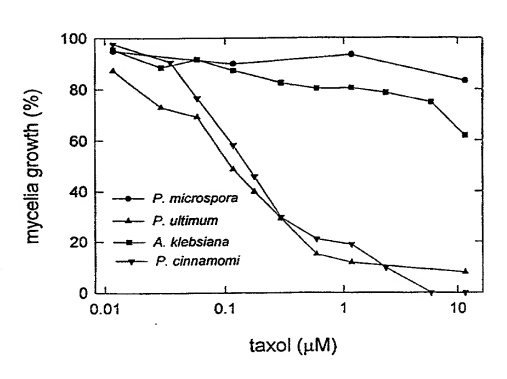


Fig. 1

CCGTCGAGCTCTACTCCAAAGAAGGCCTCTTTGTTGCTTCCTCTCAGCCTCGACATCTTCTACAAACCGCCATCATGCGTGAGATTGTT

06	CACCTCCAGACCGGTCAATGCGAAATTGGTGCTGCCTTCTGGCAACCATCTCTGGCGAGCACGGTCTCGACAGCAATGGAGTC ${ m H}$ ${ m L}$ O ${ m T}$ G ${ m G}$ O ${ m C}$ G ${ m N}$ O ${ m I}$ G ${ m A}$ A ${ m F}$ W ${ m Q}$ T ${ m I}$ S G ${ m E}$ H G ${ m L}$ D S ${ m N}$ G ${ m V}$ 35	
180		
	YNGTSELQLERMSVYFNEASGNKYVPRA65	. •
270		
	VDLEPGTMDAVRAGPFGQLFRPDNFVFGQS 95	. 0
360	sggrgccgagcrcgaccaggrccrcgacgrrgrccgrcgcgaggccgag	
	GAGNNWAKGHYTEGAELVDQVLDVVRREAE 125	ស
450	CCTGGGTGGTGCTGCTGCTGTATGGGTACTCTGTTGATCTCCAAGATC	ļ.
	ACDCLOGFQITHSLGGGTGAGMGTLLISKI 155	2
540	SGTCGTGCCCTCCCCAAGGTCTCCGACACCCGTCGTCGAGCCCTACAACGCC	
	REEFPDRMMATESVVPSPKVSDTVVESPKVB5185	വ
630	HOUSE CATTGACAACGAGGTCICIACGACAICIGCAIGCACACACAT	ιC
720	TO TO TO THE TOTAL)
9	NETUB5	
	KLSNPSYGDINHLVSAVMSGVTTCLRFPGQ 245	15
810	STGCCCTTCCTCGT(
	INSDLRKIAVNMVPPPRLHFFMVGFAPLTS 275	75
006	CGTGGCGCCCACTCTTTCCGTGCCGTCACCGTCCCCGAGTTGACCCAGCAGATGTTCGACCCCAAGAACATGATGGCTGCTTCCGACTTC	
	L T Q Q M F D)5
066	CGTAACGGTCGCTACCTGACCTGCTCTGCCATCTTCCGTGGTAAGGTCTCCATGAAGGAGGTCGAGGACCAGATGCGCAACGTCCAGAAC	
	RNGRYLTCSAIFRGKVSMKEVEDQMRNVQN 335	35
1080	AAGAACTCGTCCTACTTCGTCGAGTGGATCCCCAACAACGTGCAGACCGCCCTCTGCTCCATTCCTCCCCGCGGCCTTAAGATGTCGTCC	
	KNSSYFVEWIPNNVQTALCSIPPRGLKMSS 365	55
1170	ACTITICGICGGAAACTCGACTGCTATCCAGGAGCTGTTCAAGCGCATCGGCGAGCAGTTCACTGCCATGTTCCGTCGCAAGGCTTTCTTG	
		i.
	FKRIGEQFTAMFRRKAFL 39	ري ح
1260	STICACTGAGGCCGAGTCCAACATGAACGACTTGGTCAGCGAATACCAGCAGTAC	
	FTEAESNMNDIVSEYQQY 42	ζ Ω
1350	GAGGAGCCTCTGCCCGAGGACGAGTAAACGGCTCGCTAGAGGCTACCAAAG	
	7	4 0
1440 1530		
1620	TTAGATATACCTTTTGGGGCTGAATAAAAAAAAAAAAAA	

	30	60	06		120	(1	150	180	210	240		210 270	10	300) }	330	360		390		44 0.00	446			
	E L V H I Q G G Q C G N Q I G A K F W E V I S D E H G V CGGGTAGCTACCATGGTGACTCCGACTGGAGCGCATCAACGTGTACTACAACGAAGCTACGGGCGGTCGTTACGTG			GTCTTCGGCCAACCCGGTGCTGGTAACAACTGGGCCAAGGGTCACTATACGGAAGGCGCTGAATTGATCGACTCGGTCTTGGATGTCGCC	L I D S V L D V A	TTCCAGATCACCCACTCCCTCGGTGGTGGTACCGGTTCCGGTATGGGTACGCTT	ж A к S C D C L Q G F Q I T H S L G G G T G S G M G T L ICTAAGATCCGTGAAGAATACCCAGATCG <u>TATCATGTGCACGTACTCGGTGTGC</u> CCATCGCCAAAGGTGTCGGATACCGTCGTC	ATC		IGCITICCGIACCCIGAAGTIGACGACCCCAACGTACGGTGACTTGGTGTGTGTGCTGCTGCCATGTCCGGTATCACGACGTGCCTG CFRTLKIGAAGTIGACGACCCCAACGTACGGTGACTTGAACCACTTGGTGTGTGCTGCCATGTCCGGTATCACGACGTGCCTG	LTC	F W H	GCGCCATTGACCTCCCGTGGTTCGCA <u>GCAGTACCGTGCTTTGACCGTCCCAG</u> AATTGACCCAGCAACAATTCGACGCGAAGAACATGATG		GTCGTTATTTAACTGCTGCCTGTATGTTCCGTGGCCGCATGAGCACCAAGGAAGTCGATGAACAAATG	w	ი ლ	1	æ	PAGGCTTTCTTGCACTGGTACACGGTGAAGGTATGGATGAATTTCACGGAAGCCGAGTCGAACATGAACGATTTGGTGTCG	ккавт на матристра и правод правод В правод пра	EGEFDEDEEMDEMM*	GCGATATAGCGACTCCTTTGCAGCGGCGGTTGTGGCGGCGTCGAGATATTCTCCAAGTACCATACAGAACGTGTAGTGGACTCTTCGTAT	 TCAACTATTACTCC <u>AATATT</u> AGCGAGGTAGCTTCACTACGAGCAGGCGAGTTAGTCGCTTCCGTTCTGCTCCTACTGGAAGAGAGAG	
1	182	1	272	362		452	542	632	0	77/	812		902		992	1082		1172	(1262	1352		1442	1532	

CTGGGACGGTCAAAGCACGGTACTGCCAGCCACCGCCGCCGCCGCAGAAGCACCCCGAGTTCATTCGCAGCAAACAGACATA <u>ATG</u>AGAGAACTAGTTCACATCCAAGGTGGCCAGTGCGGTAACCAAATTGGCGCCAAGTTTTGGAAGTGATTTCTGATGAACACGGTGTG

1 92

-	###	
	CAGCGACAACATGAGAGCTCGTTCACATCCAGGGTGGCCAGTGCGGTAACCAGATCGGCGCCCAAGTTCTGGGAGGTCGTTCTCCGACGAGAG	
92	Q C G N Q I G A K F W E V V S D E ACCTGCAGCTGGAGCGCGGCGGC	27
182	OLO LE RINVYY NE AT GG	57
	S I M D S V R A G P Y G Q L F R P	87
	SGGCCARGGGACTCTACGGAGGCGCCGAGCICAICGACICGGGGGCGCGCGCGCG	117
	OGFO ITHSLGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	147
542	ORIM CTYS VCPSPRED SPKVSD AGCTTGTCGAGAACGC <u>CGATGAGGTCATGTGCCTGGATAA</u> CGAGGCCCTG PCBTUB2U	177
632	OLVENADEVMCCLDNEALC	207
	TAAGCTTGCCGTGAACCTGATCCCGCTCTGCACTTCTTCATG	237
	R L A V N L I P F P R L H F F M	267
902	TRALLTVPEELTOQQOFDAFIOACIOCOCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	297
992	I T A A C M F R G R M S T K E V D CGTCGAGTGGATCCCCAACATCAAGGC <u>T</u> AGCGTGTGACATCCCG	327
1082	E O M L N V O N K N S S Y F V E W I P N N I K A S V C D I P 3 CCTAAGGGTCTCAAGGATGTCCGAACAGTTCAGGCTT	357
172	O I A I Q E M F K K V S E Q F I A GGGTATGGATGAGAGTTCACGGAGGCTGAGTCCAACATGAACGAT	38./
1262	I G M D E M E F T E A E S N M N D GGAGGAGGCGAGTTCGACGAGGAATGGATGAGATAGACGCG (135	417 0)
	PCBTU	•
	7 Y O X O O X O D Y J Y E E E E E E E E E E E E E E E E E	44

Fig. 4

	LOXeT	binding	region	-		
.UBB-pc 122050	- • •	GOCGNOIGAKFWEVV	SDEHGVDPTGSYI	GDSDLQLERINVYY	GAKFWEVVSDEHGVDPTGSYHGDSDLQLERINVYYNEATGGRYVPRAILMDLEP I	70
.UBB-pc r22050	GTMDSVRAGI	PYGQLFRPDN <i>F</i> VFGQ'	Igagnnwakghyi	EGAELIDSVLDVVR	UBB-pc GTMDSVRAGPYGQLFRPDNFVFGQTGAGNNWAKGHYTEGAELIDSVLDVVRKEAESCDCLQGFQITHSLG 22050	140
UBB-pc 122050		LLISKIREEYPDRIM	CTYSVCPSPKVSI	TVVEPYNATLSVHQ	GGTGSGMGTLLISKIREEYPDRIMCTYSVCPSPKVSDTVVEPYNATLSVHQLVENADEVMCLDNEALYDI	210
UBB-pc 122050		Taxol binding region II CFRTLKLTTPTYGDLNHLVCAAMS N	GITTCLRFPGQL	ISDLRKLAVNLIPFP V KLF	Taxol binding region II CFRTLKLTTPTYGDLNHLVCAAMSGITTCLRFPGQLNSDLRKLAVNLIPFPRLHFFMIGFAPLTSRGSQQ V KLF	280
TUBB-pc 122050	YRALTVPEL	tqqqfdaknmmcaad	PRHGRYLTAACM	RGRMSTKEVDEQML	'UBB-pc yraltvpeltooofdaknmmcaadprhgryltaacmfrgrmstkevdeomlnvonknssyfvewipnnik 350 122050	350
.UBB-pc 122050	ASVCDIPPK(Q	GLKMSTTFIGNSTAL	Qemfkrvseqfty	MFRRKAFLHWYTGE	UBB-pc ASVCDIPPKGLKMSTTFIGNSTAIQEMFKRVSEQFTAMFRRKAFLHWYTGEGMDEMEFTEAESNMNDLVS 122050 Q	420
UBB-pc 722050	EYQQYQDAT? G	EYQQYQDATAEEEGEFDEDEEWMR G				444

					c																	
	P. microspora	spora	Н	MREIV	/HI	Ŏ.	MREIVHLOTGOCGNOIGAAFWQTISGEHGLDSNGVYNGTSELQLERMSVYFNEASGNKYV	GAAF	WOTIE	3GE	IGLD	SNG	Ž	NG	TS	ELQ	JERMS'	VYF	NEA	SGNKYV	9	
	N. Crassa					t	:					AS					Z				09	
		200	Н									GS				А	Z				9	
			l 1	H	۲.	E G		×	ΕV	Д	>	Ęď	ß	Ħ	Д	Д	H	≯ 1		T GR	9	
	P. cinnamomi	nomi	H	H				×	EW	А	>	Гď	Ŋ	I	Ω	Д	IIN	•		T GR	9	
		iana	Н	H	٦,	E E		×	ΕV	А	>	ΡŢ	ທ	H	Ω	А	H		> 1	T - T	50 0	
	Human 82		-			Ø		×	思く	Д	н	Ld	<u>F</u> -	耳	Ω	Д	NI			T G	9	
	p. microspora	spora	75	PRAVLVDLE		₹PG.	PGTMDAVRAGPFGQLFRPDNFVFGQSGAGNNWAKGHYTEGAELVDQVLDVVRREAE	GPFG	QLFR1	NOG	₹VFG	SSĞ	JAG.	Ŕ	IWA	KGH	YTEGA	EL	7DQV.	LDVVRR	EAE 1	25
	מש מ		3						ţ													125
		12 12 13					ט		M										z	^	∺	125
	P. v1timum	E .		Σ			ស					Д						-	S	A K		125
		nom 1					ល	×				H						•-	E S	L K		125
		iana					တ	≯				H						,-,	ន	×		124
	Human 82						ស		н										Ø	×		25
_														7	*	•-						
_	p. microspora	spora	AC	DCLO	GF(TIC	ACDCLOGFOITHSLGGGTGAGMGTLLISKIREEFPDRMMATFSVVPSPKVSDTVVEPYNATLSVH	GAGM	GTLL	ISK	IREE	TEL	瓦	VIMI2	\TF	SW	PSPKV	SD.	TVVE	PYNATI		190
	N. Crassa	4 (1	ט	₹		ì																1.90
		in the second	U																		-	190
			r.					ໝ				×	r1	н	بر ن						_	190
		, #C#	ı C.					ល				×	-1		C Y	υ 					71	190
	r. Clobetons) V.					ı vo				þ	-								_	189
	A. Aleba. Human β2	5	ນ			Н		S				≯	.7	Н	z						H	06.
					*	*		Ħ	Taxol binding region	hind	ing	cegi	g	II	H		•-	*				
	P. microspora	spora	OI	VENS	DE	TFC	OLVENSDETFCIDNEALYDICMRTLKLSNPSYGDLNHLVSAVMSGVTTCLRFPGQLNSDLRKLAV	DICE	RTLK	LSN	PSYC	TOT.	日	LV	SAT	MSG	VTTCL	RF	PGQL	NSDLRK	TLAV 2	522
	N. Crassa	, u	ł														VS					255
		200		Ħ																	W 2	522
		5		Æ		M.	ı	ĒΨ	r.	TI	H			_		Ą	н				.,	255
		momi		æ		M	7	E 4	F	TT	H			_		ď.	н				.,	255
		iana		ø		MA	1	ĮΣţ	F-	H	<u>-</u> -				ري ت	Æ	н			>	.,	253
	Human 82			H		×		ĮΞŧ	r-	II	H				-	E				Ą	• • •	255

Fig. 6A

				#=	‡												
P. microspora		PRLH	FFM	/GFA	PLTSR	GAHSFR	NMVPFPRLHFFWVGFAPLTSRGAHSFRAVTVPELTQQMFDPKNMMAASDFRNGRYLTCSAIFRGK	ZOMF	DPK	NIMIMA	ASI	OFF	NGRY	TLTCS	LFRG		320
N. crassa						Ħ	ល									m	20
A. nidulans						×	Ø									(L)	20
P. ultimum	LI		H			SQQY	ŗ	O	ď	ບ	4	а	H	AACM		R 3	320
P. cinnamomi	ΓI		H			SQQY	디	Ø	Ą	ບ	ď	а	田	AACM		R 3	20
A. klebsiana	ΓΊ		Н	_		SQQY	니	Ø	ď	บ	Ø	Д	H	AACM		R 3	18
Human \$2			 1	Ωı		SOOY	ij		ø		บ	Д	Ħ	VA	>	3	20
										#							
P. microspora	VSMKE	VEDQ	MRN	/QNK	NSSYF	VEWIPN	VSMKEVEDQMRNVQNKNSSYFVEWIPNNVQTALCSIPPRGLKMSSTFVGNSTAIQELFKRIGEQF	IPPR	GLK	MSST	FV	SKE	TAIO	ELFKF	LGEO		385
N. crassa																ന	385
A. nidulans			11	H S	Ø		N S				H		ഗ		Q V	m	385
P. ultimum	Η	DE	ы				IKASV D	×		H				Σ	ΛS	m	385
P. cinnamomi	Ε	DE	ᆸ				IKASV D	×		H	H			Z	VS	m	385
 A. klebsiana	H M	DE	Ы				IKASV D	X		H	H			Σ	ΛS	ćζ	383
 Human β2	×	DE	7				K V D			Ø	Н				လ	M	85
 P. microspora	TAMFF	RKAF	LHW	TGE	GMDEM	EFTEAE	TAMFRRKAFLHWYTGEGMDEMEFTEAESNWNDLVSEYQQYQDAGVDEEEEEYEEEP-LPEDE\$	EYQC	YQD.	AGVD	田田田	国国	YEEE	P-LPE	3DE\$	4	446
N. crassa														A-PL	G E\$	4,	447
A. nidulans										SIS	O		ø	E-IM	G EŞ	4	447
P. ultimum										TAE	1	ט	FD	реемр мм\$) MM\$	4	446
P. cinnamomi										TAE	1	ש	ED FD	DEEWMR\$	R\$	4	444
A. klebsiana										TAE	I	ರ	ED ED	DEEMD MM\$) MM\$	4	444
Human 82										TAE	1	ש	阳	AEEEVA\$:VA\$	4	445

Fig. 6B

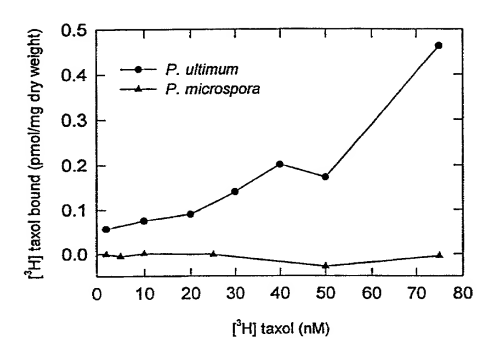


Fig. 7A

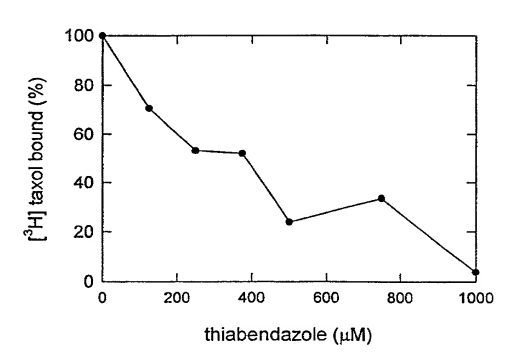


Fig.7B

	•	Region	Region I (I-31 aa)	1-31 00) ********	***	*		Regio	****** 7) II 0	Region II (212-231 aa *****	aa)
(s) Pig	MREIV	HIQAG	MREIVHIQAGQCGNQIGAKFWEVISDEHGID	GAKF	WEVI	SDEH	GID	FRTL	KLTTPTY	FRTLKLTTPTYGDLMHLVSA	/SA
(s) Human β 2		ដ									
(s) Drosophila β1					H						H
(s) Xenopus $\beta 4$		H									
(s) Tetrahymena			_O								
(s) Physarum β_1											
(s) P. ultimum	ы	Ö					>				ບ
(s) P. cinnamomi	H	ರ			>		>				ບ
(r)A. klebsiana	ы	ტ					>		Z		ບ
(r) P. microspora		I.		ď	ğ	ტ	니	Σ	SNS		
(r)A. nidulans benA		T I		ø	ď	ש	ᆸ	Z	SNS		
(r) S. cerevisiae	H	SA	≯	ø	H	ဗ္ဗ	ᄓ	œ	SÖN	Z	လ

Fig. 8

DECLARATION AND POWER OF ATTORNEY



As a below-named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name, that I believe that I am the original, first and sole inventor (if only one name is listed below) or I believe that we are the original, first and joint inventors (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention, design or discovery entitled FUNGAL BETA-TUBULIN GENES, the specification of which (check one)

- () is attached hereto; or
- (X) was filed on March 23, 2000, as PCT International Application Number PCT/US00/07995, and amended by letter to the International Preliminary Examining Authority received on May 21, 2001

that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above; that I do not know and do not believe that said invention, design or discovery was ever known or used in the United States of America before my invention or discovery thereof, or patented or described in any printed publication in any country before my invention or discovery thereof, or more than one year prior to this application, or in public use or on sale in the United States of America more than one year prior to this application; that said invention, design or discovery has not been patented or made the subject of an inventor's certificate issued prior to the date of this application in any country foreign to the United States of America on an application filed by me or my legal representatives or assigns; and that I acknowledge the duty to disclose information of which I am aware which is material to the examiner of this application in accordance with 37 C.F.R. § 1.56(a).

I hereby claim foreign priority benefits under 35 U.S.C. § 119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application(s) for patent or inventor's certificate having a filing date before that of the application on which priority is claimed:

NUMBER COUNTRY

DATE FILED

PRIORITY CLAIMED

N/A

I hereby claim the benefit under Title 35, United States Code, § 119(e) of any United States provisional application(s) listed below.

APPLICATION SERIAL NO.

DATE FILED

60/125,717

March 23, 1999

I hereby claim the benefit under 35 U.S.C. § 120 of any United States Application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States Application(s) in the manner provided by the first paragraph of 35 U.S.C. § 112, I acknowledge the duty to disclose material information as defined in 37 C.F.R. § 1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application:

APPLICATION SERIAL NO.

DATE FILED

STATUS

N/A

I hereby appoint:

3

Eugenia S. Hansen Karen L. Knezek Rod A. Cooper Reg. No. 31,966, Reg. No. 39,253, Reg. No. 42,436

all of the firm of Sidley & Austin, my attorneys with full power of substitution and revocation, to prosecute this application and to transact all business in the United States Patent and Trademark Office connected therewith, and to file and prosecute any international patent applications filed thereon before any international authorities under the Patent Cooperation Treaty.

Send correspondence to:

Direct telephone calls to:

Sidley Austin Brown & Wood 717 N. Harwood Suite 3400

Dallas, TX 75201-6507

Eugenia S. Hansen at (214) 981-3300

Atty. Docket No. 10365/07304

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under § 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

DECLARATION AND POWER OF ATTORNEY

As a below-named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name, that I believe that I am the original, first and sole inventor (if only one name is listed below) or I believe that we are the original, first and joint inventors (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention, design or discovery entitled FUNGAL BETA-TUBULIN GENES, the specification of which (check one)

- () is attached hereto; or
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that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above; that I do not know and do not believe that said invention, design or discovery was ever known or used in the United States of America before my invention or discovery thereof, or patented or described in any printed publication in any country before my invention or discovery thereof, or more than one year prior to this application, or in public use or on sale in the United States of America more than one year prior to this application; that said invention, design or discovery has not been patented or made the subject of an inventor's certificate issued prior to the date of this application in any country foreign to the United States of America on an application filed by me or my legal representatives or assigns; and that I acknowledge the duty to disclose information of which I am aware which is material to the examiner of this application in accordance with 37 C.F.R. § 1.56(a).

I hereby claim foreign priority benefits under 35 U.S.C. § 119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application(s) for patent or inventor's certificate having a filing date before that of the application on which priority is claimed:

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APPLICATION SERIAL NO.

DATE FILED

STATUS

N/A

I hereby appoint:



Eugenia S. Hansen Karen L. Knezek Rod A. Cooper

Reg. No. 31,966 Reg. No. 39,253 Reg. No. 42,436

all of the firm of Sidley & Austin, my attorneys with full power of substitution and revocation, to prosecute this application and to transact all business in the United States Patent and Trademark Office connected therewith, and to file and prosecute any international patent applications filed thereon before any international authorities under the Patent Cooperation Treaty.

Send correspondence to:

Direct telephone calls to:

Sidley Austin Brown & Wood 717 N. Harwood Suite 3400 Dallas, TX 75201-6507 Eugenia S. Hansen at (214) 981-3300

Atty. Docket No. 10365/07304

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under § 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Inventor's signature: <u>Kajinder Sidly</u> Date: 10 - 26 - 0Residence (City, County, State): Dallas, Dallas County, Texas Citizenship: United States of America Post Office Address: 7521 Danfield Court, Dallas, Texas 75252 Full name of sole or joint inventor: Arthur P. Bollon Inventor's signature: Residence (City, County, State): Dallas, Dallas County, Texas Citizenship: United States of America Post Office Address: 13227 Cedar Lane, Dallas, Texas 75234 Full name of sole or joint inventor: Jing-Hong Mu Inventor's signature: Date: Residence (City, County, State): Dallas, Dallas County, Texas

Post Office Address: 534 Newberry Drive, Richardson, Texas 75080

Full name of sole or joint inventor: Rajinder S. Sidhu

Citizenship: China

*	Full name of sole or joint inventor: Rajinder S. Sidnu
M	Inventor's signature: Lajinder 5-64 Sidler
	Date: 10-26-01
	Residence (City, County, State): Dallas, Dallas County, Texas
	Citizenship: United States of America
	Post Office Address: 7521 Danfield Court, Dallas, Texas 75252
1 2 2	
- (1)	Full name of sole or joint inventor: Arthur P. Bollon
9W	Inventor's signature: Autof Bullet
Ź	Date: october 12, 2001
	Residence (City, County, State): Dallas, Dallas County, Texas
The work	Citizenship: United States of America
#	Post Office Address: 13227 Cedar Lane, Dallas, Texas 75234
A STATE OF THE STA	
	Full name of sole or joint inventor: Jing-Hong Mu
	Inventor's signature: January Mari
V	Date: 01, 12, 00
	Residence (City, County, State): Dallas, Dallas County, Texas

Post Office Address: 534 Newberry Drive, Richardson, Texas 75080

Full name of sole or joint inventor: Rajinder S. Sidhu

Citizenship: China

Applicant or Patentee:

Rajinder S. Sidhu, et al.

Serial or Patent No.:

To be assigned

International Appln. No.: International Filing Date: PCT/US00/07995 March 23, 2000

For:

FUNGAL BETA-TUBULIN GENES

VERIFIED STATEMENT (DECLARATION)

CLAIMING SMALL ENTITY STATUS

(37 C.F.R. §§ 1.9(f) AND 1.27(C)) SMALL BUSINESS CONCERN

I hereby declare that I am an official of the small business concern empowered to act on behalf of the concern identified below:

NAME OF ORGANIZATION:

Cytoclonal Pharmaceutics, Inc.

ADDRESS OF ORGANIZATION: 2110 Research Row, Suite 621, Dallas, Texas 75235

I hereby declare that the above-identified small business concern qualifies as a small business concern as defined in 13 C.F.R. § 121.12, and reproduced in 37 C.F.R. § 1.9(d), for purposes of paying reduced fees under §§ 41(a) and (b) of Title 35, United States Code, in that the number of employees of the concern, including those of its affiliates, does not exceed 500 persons. For purposes of this statement, (1) the number of employees of the business concern is the average over the previous fiscal year of the concern of the persons employed on a fulltime, part-time or temporary basis during each of the pay periods of the fiscal year, and (2) concerns are affiliates of each other when either, directly or indirectly, one concern controls or has the power to control the other, or a third party or parties controls or has the power to control both.

I hereby declare that rights under contracts or law have been conveyed to and remain with the small business concern identified above with regard to the invention, entitled FUNGAL BETA-TUBULIN GENES by inventor Rajinder S. Sidhu, et al.

described in:

- () the specification filed herewith.
- International Application No. PCT/US00/07995, filed March 23, 2000 and (X) amended by letter to the International Preliminary Examining Authority received on May 21, 2001.
- () patent no., issued.

If the rights held by the above-identified small business concern are not exclusive, each individual, concern or organization having rights to the invention is listed below* and no rights to the invention are held by any person, other than the inventor, who could not qualify as an independent inventor under 37 C.F.R. § 1.9(c) if that person made the invention or by any

VERIFIED STATEMENT (DECLARATION) **CLAIMING SMALL ENTITY STATUS** INTERNATIONAL PATENT APPLICATION NO. PCT/US00/07995 PAGE 1 OF 2

Applicant or Patentee:

Rajinder S. Sidhu, et al.

Serial or Patent No.:

To be assigned

International Appln. No.: International Filing Date: PCT/US00/07995 March 23, 2000

For:

FUNGAL BETA-TUBULIN GENES



VERIFIED STATEMENT (DECLARATION)

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I hereby declare that rights under contracts or law have been conveyed to and remain with the small business concern identified above with regard to the invention, entitled FUNGAL BETA-TUBULIN GENES by inventor Rajinder S. Sidhu, et al.

described in:

- the specification filed herewith. ()
- International Application No. PCT/US00/07995, filed March 23, 2000 and (X) amended by letter to the International Preliminary Examining Authority received on May 21, 2001.
- patent no., issued. ()

If the rights held by the above-identified small business concern are not exclusive, each individual, concern or organization having rights to the invention is listed below* and no rights to the invention are held by any person, other than the inventor, who could not qualify as an independent inventor under 37 C.F.R. § 1.9(c) if that person made the invention or by any

VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS INTERNATIONAL PATENT APPLICATION NO. PCT/US00/07995 PAGE 1 OF 2

concern which would not qualify as a small business concern under 37 C.F.R. § 1.9(d), or a nonprofit organization under 37 C.F.R. § 1.9(e).

Each person, concern or organization having any rights in the invention is listed below:

- no such person, concern or organization (X)
- persons, concerns or organizations listed below* ()

*NOTE: Separate verified statements are required from each named person, concern, or organization having rights to the invention averring to their status as small entities. (37 C.F.R. § 1.27)

NAME:

ADDRESS:

nonprofit organization small business concern () () individual

I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate. (37 C.F.R. § 1.28(b))

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under § 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.

NAME OF PERSON SIGNING:

Robert J. Rousseau

TITLE IN ORGANIZATION:

Vice President of Business Development and Licensing,

Cytoclonal Pharmaceutics, Inc.

ADDRESS OF PERSON SIGNING:

2110 Research Row, Suite 621, Dallas, Texas

75235

SIGNATURE: Robert J. Rousseaw

DATE: 10/10/01

concern which would not qualify as a small business concern under 37 C.F.R. § 1.9(d), or a nonprofit organization under 37 C.F.R. § 1.9(e).

Each person, concern or organization having any rights in the invention is listed below:

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- persons, concerns or organizations listed below* ()

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ADDRESS:

nonprofit organization small business concern individual ()

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SIGNATURE: Robert J. Rousseau DATE: 10/10/01

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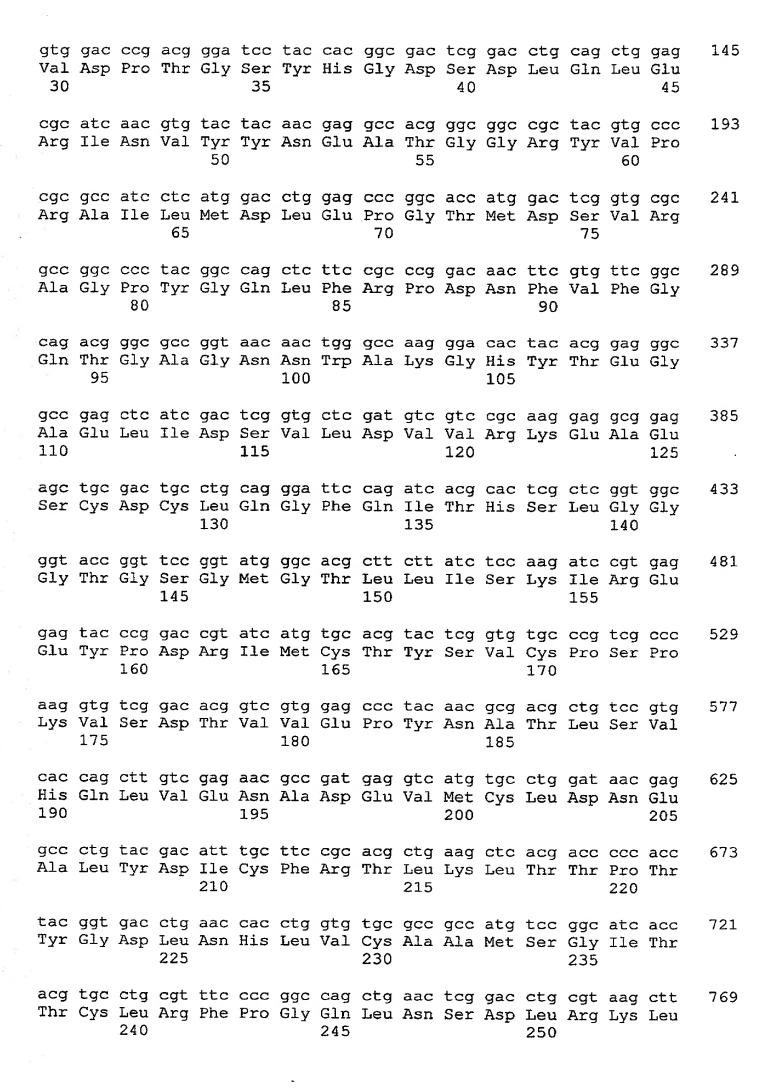
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Mr. Com Mark

351

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245

20

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Ser Gly Val Tyr Asn Gly Thr Ser Glu Leu Gln Leu Glu Arg Met Asn 35 40 45

Val Tyr Phe Asn Glu Ala Ser Gly Asn Lys Tyr Val Pro Arg Ala Val 50 55 60 Leu Val Asp Leu Glu Pro Gly Thr Met Asp Ala Val Arg Ala Gly Pro Phe Gly Gln Leu Phe Arg Pro Asp Asn Phe Val Phe Gly Gln Ser Gly Ala Gly Asn Asn Trp Ala Lys Gly His Tyr Thr Glu Gly Ala Glu Leu Val Asp Gln Val Leu Asp Val Val Arg Arg Glu Ala Glu Gly Cys Asp Cys Leu Gln Gly Phe Gln Ile Thr His Ser Leu Gly Gly Gly Thr Gly 135 Ala Gly Met Gly Thr Leu Leu Ile Ser Lys Ile Arg Glu Glu Phe Pro Asp Arg Met Met Ala Thr Phe Ser Val Val Pro Ser Pro Lys Val Ser Asp Thr Val Val Glu Pro Tyr Asn Ala Thr Leu Ser Val His Gln Leu Val Glu Asn Ser Asp Glu Thr Phe Cys Ile Asp Asn Glu Ala Leu Tyr Asp Ile Cys Met Arg Thr Leu Lys Leu Ser Asn Pro Ser Tyr Gly Asp Leu Asn His Leu Val Ser Ala Val Met Ser Gly Val Thr Val Ser Leu Arg Phe Pro Gly Gln Leu Asn Ser Asp Leu Arg Lys Leu Ala Val Asn Met Val Pro Phe Pro Arg Leu His Phe Phe Met Val Gly Phe Ala Pro 260 Leu Thr Ser Arg Gly Ala His His Phe Arg Ala Val Ser Val Pro Glu 280 Leu Thr Gln Gln Met Phe Asp Pro Lys Asn Met Met Ala Ala Ser Asp Phe Arg Asn Gly Arg Tyr Leu Thr Cys Ser Ala Ile Phe Arg Gly Lys 315 Val Ser Met Lys Glu Val Glu Asp Gln Met Arg Asn Val Gln Asn Lys Asn Ser Ser Tyr Phe Val Glu Trp Ile Pro Asn Asn Val Gln Thr Ala Leu Cys Ser Ile Pro Pro Arg Gly Leu Lys Met Ser Ser Thr Phe Val 355 360 365

Gly Asn Ser Thr Ala Ile Gln Glu Leu Phe Lys Arg Ile Gly Glu Gln 370 375 380

Phe Thr Ala Met Phe Arg Arg Lys Ala Phe Leu His Trp Tyr Thr Gly 385 390 395 400

Glu Gly Met Asp Glu Met Glu Phe Thr Glu Ala Glu Ser Asn Met Asn 405 410 415

Asp Leu Val Ser Glu Tyr Gln Gln Tyr Gln Asp Ala Gly Val Asp Glu 420 425 430

Glu Glu Glu Glu Glu Glu Glu Ala Pro Leu Glu Glu Glu 435 440 445

<210> 26

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Gly Ala Ala Phe Trp Gln Thr Ile Ser Gly Glu His Gly Leu Asp Gly 20 25 30

Ser Gly Val Tyr Asn Gly Thr Ser Asp Leu Gln Leu Glu Arg Met Asn 35 40 45

Val Tyr Phe Asn Glu Ala Ser Gly Asn Lys Tyr Val Pro Arg Ala Val 50 55 60

Leu Val Asp Leu Glu Pro Gly Thr Met Asp Cys Val Arg Ala Gly Pro 65 70 75 80

Phe Gly Glu Leu Phe Arg Pro Asp Asn Phe Val Phe Gly Gln Ser Gly 85 90 95

Ala Gly Asn Asn Trp Ala Lys Gly His Tyr Thr Glu Gly Ala Glu Leu 100 105 110

Val Asp Asn Val Val Asp Val Val Arg Arg Glu Ala Glu Gly Cys Asp 115 120 125

Cys Leu Gln Gly Phe Gln Ile Thr His Ser Leu Gly Gly Gly Thr Gly 130 135 140

Ala Gly Met Gly Thr Leu Leu Ile Ser Lys Ile Arg Glu Glu Phe Pro 145 150 155 160 Asp Arg Met Met Ala Thr Phe Ser Val Val Pro Ser Pro Lys Val Ser Asp Thr Val Val Glu Pro Tyr Asn Ala Thr Leu Ser Val His Gln Leu Val Glu His Ser Asp Glu Thr Phe Cys Ile Asp Asn Glu Ala Leu Tyr 200 Asp Ile Cys Met Arg Thr Leu Lys Leu Ser Asn Pro Ser Tyr Gly Asp Leu Asn His Leu Val Ser Ala Val Met Ser Gly Val Thr Thr Cys Leu 230 Arg Phe Pro Gly Gln Leu Asn Ser Asp Leu Arg Lys Trp Ala Val Asn 245 250 Met Val Pro Phe Pro Arg Leu His Phe Phe Met Val Gly Phe Ala Pro 265 Leu Thr Ser Arg Gly Ala Tyr Ser Phe Arg Ala Val Ser Val Pro Glu 280 Leu Thr Gln Gln Met Phe Asp Pro Lys Asn Met Met Ala Ala Ser Asp Phe Arg Asn Gly Arg Tyr Leu Thr Cys Ser Ala Ile Phe Arg Gly Lys Val Ser Met Lys Glu Val Glu Asp Gln Met Arg Asn Ile Gln Ser Lys Asn Gln Ser Tyr Phe Val Glu Trp Ile Pro Asn Asn Ile Gln Ser Ala 345 Leu Cys Ser Ile Pro Pro Arg Gly Leu Lys Met Ser Ser Thr Phe Ile Gly Asn Ser Thr Ser Ile Gln Glu Leu Phe Lys Arg Val Gly Asp Gln Phe Thr Ala Met Phe Arg Arg Lys Ala Phe Leu His Trp Tyr Thr Gly Glu Gly Met Asp Glu Met Glu Phe Thr Glu Ala Glu Ser Asn Met Asn Asp Leu Val Ser Glu Tyr Gln Gln Tyr Gln Asp Ala Ser Ile Ser Glu 420 425 Gly Glu Glu Glu Tyr Ala Glu Glu Glu Ile Met Glu Gly Glu Glu 440 435

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Ile Cys Phe Arg Thr Leu Lys Leu Thr Asn Pro Thr Tyr Gly Asp Leu 210 215 220

Asn His Leu Val Cys Ala Ala Met Ser Gly Ile Thr Thr Leu Leu Arg 225 230 235 240

Phe Pro Gly Gln Leu Asn Ser Val Leu Lys Leu Ala Val Asn Leu Ile 245 250 255 Pro Phe Pro Arg Leu His Phe Phe Met Ile Gly Phe Ala Pro Leu Thr 265 Ser Arg Gly Ser Gln Gln Tyr Arg Ala Leu Thr Val Pro Glu Leu Thr 280 Gln Gln Gln Phe Asp Ala Lys Asn Met Met Cys Ala Ala Asp Pro Arg His Gly Arg Tyr Leu Thr Ala Ala Cys Met Phe Arg Gly Arg Met Ser Thr Lys Glu Val Asp Glu Gln Met Leu Asn Val Gln Asn Lys Asn Ser Ser Tyr Phe Val Glu Trp Ile Pro Asn Asn Ile Lys Ala Ser Val Cys 345 Asp Ile Pro Pro Lys Gly Leu Lys Met Ser Thr Thr Phe Ile Gly Asn Ser Thr Ala Ile Gln Glu Met Phe Lys Arg Val Ser Glu Gln Phe Thr Ala Met Phe Arg Arg Lys Ala Phe Leu His Trp Tyr Thr Gly Glu Gly 395 Met Asp Glu Met Glu Phe Thr Glu Ala Glu Ser Asn Met Asn Asp Leu Val Ser Glu Tyr Gln Gln Tyr Gln Asp Ala Thr Ala Glu Glu Glu Gly 425 Glu Phe Asp Glu Asp Glu Glu Met Asp Glu Met Met

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Leu Val Ser Leu
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Leu Val Ser Ala

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Leu Val Ser Ala
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<211> 31 <212> PRT <213> Pestalotiopsis microspora

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Leu Val Ser Ala 20

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Leu Val Ser Ala 20

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Leu Val Ser Ser 20

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of:

Rajinder S. Sidhu, et al.

For:

FUNGAL BETA-TUBULIN GENES

Application Serial No.

Unassigned

Filing Date:

Concurrently herewith

International

Application No.:

PCT/US00/07995

International

Filing Date:

23 March 2000

Assistant Commissioner for Patents

Box PCT

Washington, D.C. 20231

EXPRESS MAIL NO. <u>EL794555629US</u> DATE OF DEPOSIT 19 September 2001
I hereby certify that this correspondence is being deposited with the United States Postal Service "Express Mail Post Office to Addressee" service under 37 CFR 1.10 on the date indicated above and is addressed to the Assistant Commissioner for Patents, Box PCT, Washington, D.C. 20231 on 19 September 2001
(Date of Deposit)

Derrick Gordon

Name of Depositor

Signature

Date of Signature: 19 September 2001

Dear Sir:

STATEMENT UNDER WIPO STANDARD ST.25

This statement is being filed pursuant to the WIPO Standard ST.25 and pursuant to Administrative Instructions, Section 208. Applicants are filing concurrently herewith a Sequence Listing in paper copy and computer readable form. The Sequence Listing has been amended to delete the corporate applicant CYTOCLONAL PHARMACEUTICS, INC. from the applicant list, and the Attorney Docket No. has been amended to reflect the Attorney Docket No. for the national stage application. The undersigned hereby states that the sequence information recorded in computer readable form is identical to the written Sequence Listing furnished in the international patent application as filed and that the Sequence Listing does not go beyond the international application. The undersigned hereby states that the Sequence Listing in paper copy filed concurrently herewith and the sequence listing in computer readable form are the same.

Respectfully submitted,

Karen L. Knezek

Registration No. 39,253

KLK\ld 19 September 2001 Sidley Austin Brown & Wood 717 N. Harwood, Suite 3400 Dallas, Texas 75201-6507 (214) 981-3300

JC16 Rec'd PCT/PTO SEP 1 9 2001

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Val Tyr Phe Asn Glu Ala Ser Gly Asn Lys Tyr Val Pro Arg Ala Val Leu Val Asp Leu Glu Pro Gly Thr Met Asp Ala Val Arg Ala Gly Pro Phe Gly Gln Leu Phe Arg Pro Asp Asn Phe Val Phe Gly Gln Ser Gly Ala Gly Asn Asn Trp Ala Lys Gly His Tyr Thr Glu Gly Ala Glu Leu Val Asp Gln Val Leu Asp Val Val Arg Arg Glu Ala Glu Ala Cys Asp Cys Leu Gln Gly Phe Gln Ile Thr His Ser Leu Gly Gly Gly Thr Gly Ala Gly Met Gly Thr Leu Leu Ile Ser Lys Ile Arg Glu Glu Phe Pro 155 150 Asp Arg Met Met Ala Thr Phe Ser Val Val Pro Ser Pro Lys Val Ser 170 165 Asp Thr Val Val Glu Pro Tyr Asn Ala Thr Leu Ser Val His Gln Leu 185 Val Glu Asn Ser Asp Glu Thr Phe Cys Ile Asp Asn Glu Ala Leu Tyr 200 Asp Ile Cys Met Arg Thr Leu Lys Leu Ser Asn Pro Ser Tyr Gly Asp 215 Leu Asn His Leu Val Ser Ala Val Met Ser Gly Val Thr Thr Cys Leu 230 Arg Phe Pro Gly Gln Leu Asn Ser Asp Leu Arg Lys Leu Ala Val Asn 250 Met Val Pro Phe Pro Arg Leu His Phe Phe Met Val Gly Phe Ala Pro 265 260 Leu Thr Ser Arg Gly Ala His Ser Phe Arg Ala Val Thr Val Pro Glu 280 Leu Thr Gln Gln Met Phe Asp Pro Lys Asn Met Met Ala Ala Ser Asp 290 Phe Arg Asn Gly Arg Tyr Leu Thr Cys Ser Ala Ile Phe Arg Gly Lys Val Ser Met Lys Glu Val Glu Asp Gln Met Arg Asn Val Gln Asn Lys Asn Ser Ser Tyr Phe Val Glu Trp Ile Pro Asn Asn Val Gln Thr Ala 340

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Gly	Asn 370	Ser	Thr	Ala	Ile	Gln 375	Glu	Leu	Phe	Lys	Arg 380	Ile	Gly	Glu	Gln	
Phe 385	Thr	Ala	Met	Phe	Arg 390	Arg	Lys	Ala	Phe	Leu 395	His	Trp	Tyr	Thr	Gly 400	
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gagt	ttcai	ttc (gcag	cgaaa	ac a	gaca	gacai							cac a		112
caa Gln	ggt Gly	ggc Gly 10	cag Gln	tgc Cys	ggt Gly	aac Asn	caa Gln 15	att Ile	ggc Gly	gcc Ala	aag Lys	ttt Phe 20	tgg Trp	gaa Glu	gtg Val	160
					ggt Gly											208
					gag Glu 45											256
ggc Gly	ggt Gly	cgt Arg	tac Tyr	gtg Val 60	cct Pro	cgt Arg	gcg Ala	atc Ile	ttg Leu 65	atg Met	gat Asp	ttg Leu	gag Glu	cca Pro 70	ggt Gly	304
acc Thr	atg Met	gac Asp	tcg Ser 75	gtc Val	cgt Arg	gcc Ala	ggt Gly	cca Pro 80	ttc Phe	ggt Gly	cag Gln	ctt Leu	ttc Phe 85	cgc Arg	cca Pro	352

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				_	ggt Gly	-					_		_			544
		_		_	gaa Glu •	_			_	_		_	_	_		592
	_	_			cca Pro	_	_	_			_		_			640
	_	_		_	gtc Val		_	_	_	-		_	_	_	_	688
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					acg Thr											784
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					ctc Leu		-		_		_			_		880
			_		ggt Gly				_			_		_	_	928
					acc Thr 285			_	_		_				-	976
					tgc Cys											1024

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gaa caa atg ctc aac gtg cag aac aag aac tcg tcg tac ttt gtg gag 1120 Glu Gln Met Leu Asn Val Gln Asn Lys Asn Ser Ser Tyr Phe Val Glu 330 335 340)
tgg att cca aac atc aag gcc agc gtg tgt gat atc cca cca aag 1168 Trp Ile Pro Asn Asn Ile Lys Ala Ser Val Cys Asp Ile Pro Pro Lys 345 350 355	}
ggt ctg aag atg agt acc acc ttc gtt ggt aac tcg act gcg atc cag Gly Leu Lys Met Ser Thr Thr Phe Val Gly Asn Ser Thr Ala Ile Gln 360 370 375	õ
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aag get tte ttg cac tgg tac acg ggt gaa ggt atg gat gag atg gaa 1312 Lys Ala Phe Leu His Trp Tyr Thr Gly Glu Gly Met Asp Glu Met Glu 395 400 405	2
ttc acg gaa gcc gag tcg aac atg aac gat ttg gtg tcg gaa tac cag Phe Thr Glu Ala Glu Ser Asn Met Asn Asp Leu Val Ser Glu Tyr Gln 410 415 420)
cag tac caa gac gcg aca gca gaa gag gaa ggt gaa ttc gac gaa gat 1408 Gln Tyr Gln Asp Ala Thr Ala Glu Glu Glu Glu Phe Asp Glu Asp 425 430 435	}
gaa gaa atg gac gaa atg tagacgacgc gggcgatata gcgactcctt 1459 Glu Glu Met Asp Glu Met Met 440 445	}
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aaaaaaaaa a 1650)
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	Ser	Ser	Tyr 340	Phe	Val	Glu	Trp	Ile 345	Pro	Asn	Asn	Ile	Lys 350	Ala	Ser	
Val	Суѕ	Asp 355	Ile	Pro	Pro	Lys	Gly 360	Leu	Lys	Met	Ser	Thr 365	Thr	Phe	Val	
Gly	Asn 370	Ser	Thr	Ala	Ile	Gln 375	Glu	Met	Phe	Lys	Arg 380	Val	Ser	Glu	Gln	
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Glu	Gly	Met	Asp	Glu 405	Met	Glu	Phe	Thr	Glu 410	Ala	Glu	Ser	Asn	Met 415	Asn	
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		-														
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	acg Thr 95															337
_	gag Glu			_	_			_	_	_	_	_				385
	tgc Cys															433
	acc Thr						_							_		481
	tac Tyr	_	_	_		-	-	_		_		_	-	_		529
	gtg Val 175						-				_	-	_			577
	cag Gln		_	_		-	-		_	_	_	_	_			625
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	tgc Cys															769
	gtg Val 255															817
	gcc Ala		-					_	_			_	_	_	_	865
	ccc Pro															913
_	gct Ala	-		_			_				_		_	_		961

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gca Ala 430	gag Glu	gag Glu	gag Glu	ggc Gly	gag Glu 435	ttc Phe	gac Asp	gag Glu	gac Asp	gag Glu 440	gaa Glu	tgg Trp	atg Met	aga Arg		1342
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Gly	Ala	Lys	Phe 20		Glu	Val	Val	Ser 25		Glu	His	Gly	Val 30		Pro	
Thr	Gly	Ser 35		His	Gly	Asp	Ser 40		Leu	Gln	Leu	Glu 45	Arg	Ile	Asn	
Val	Tyr 50		Asn	. Glu	Ala	Thr		· Gly	Arg	Tyr	Val 60		Arg	Ala	Ile	
Leu 65		Asp	Leu	Glu	Pro		Thr	Met	. Asp	Ser 75		Arg	Ala	. Gly	Pro 80	

Tyr	Gly	Gln	Leu	Phe 85	Arg	Pro	Asp	Asn	Phe 90	Val	Phe	Gly	Gln	Thr 95	Gly
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Ile	Asp	Ser 115	Val	Leu	Asp	Val	Val 120	Arg	Lys	Glu	Ala	Glu 125	Ser	Cys	Asp
Cys	Leu 130	Gln	Gly	Phe	Gln	Ile 135	Thr	His	Ser	Leu	Gly 140	Gly	Gly	Thr	Gly
Ser 145	Gly	Met	Gly	Thr	Leu 150	Leu	Ile	Ser	Lys	Ile 155	Arg	Glu	Glu	Tyr	Pro 160
Asp	Arg	Ile	Met	Cys 165	Thr	Tyr	Ser	Val	Cys 170	Pro	Ser	Pro	Lys	Val 175	Ser
Asp	Thr	Val	Val 180	Glu	Pro	Tyr	Asn	Ala 185	Thr	Leu	Ser	Val	His 190	Gln	Leu
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Leu 225	Asn	His	Leu	Val	Cys 230	Ala	Ala	Met	Ser	Gly 235	Ile	Thr	Thr	Cys	Leu 240
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			Phe 260					265					270		
		275	Arg				280					285			
	290		Gln			295					300				
305			Gly		310					315					320
			Lys	325					330					335	
			Tyr 340					345					350		
		355					360	l				365	i		
Gly	Asn 370		Thr	Ala	Ile	Gln 375		Met	. Phe	e Lys	380		. Ser	: Glu	Gln

Phe Thr Ala Met Phe Arg Arg Lys Ala Phe Leu His Cys Thr Arg Gly

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390
385
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Tyr	Gly	Gln	Leu	Phe 85	Arg	Pro	Asp	Asn	Phe 90	Val	Phe	Gly	Gln	Thr 95	Gly
Ala	Gly	Asn	Asn 100	Trp	Ala	Lys	Gly	His 105	Tyr	Thr	Glu	Gly	Ala 110	Glu	Leu
Ile	Asp	Ser 115	Val	Leu	Asp	Val	Val 120	Arg	Lys	Glu	Ala	Glu 125	Ser	Cys	Asp
Cys	Leu 130	Gln	Gly	Phe	Gln	Ile 135	Thr	His	Ser	Leu	Gly 140	Gly	Gly	Thr	Gly
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Asp	Arg	Ile	Met	Cys 165	Thr	Tyr	Ser	Val	Cys 170	Pro	Ser	Pro	Lys	Val 175	Ser
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Val Cys Asp Ile Pro Pro Gln Gly Leu Lys Met Ser Thr Thr Phe Ile 355 360 365

Gly Asn Ser Thr Ala Ile Gln Glu Met Phe Lys Arg Val Ser Glu Gln 370 375 380

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Glu Gly Met Asp Glu Met Glu Phe Thr Glu Ala Glu Ser Asn Met Asn 405 410 415

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Val Tyr Tyr Asn Glu Ala Thr Gly Gly Lys Tyr Val Pro Arg Ala Val 50 55 60

Leu Val Asp Leu Glu Pro Gly Thr Met Asp Ser Val Arg Ser Gly Pro 65 70 75 80

Phe Gly Gln Ile Phe Arg Pro Asp Asn Phe Val Phe Gly Gln Ser Gly 85 90 95

Ala Gly Asn Asn Trp Ala Lys Gly His Tyr Thr Glu Gly Ala Glu Leu 100 105 110

Val Asp Ser Val Leu Asp Val Val Arg Lys Glu Ala Glu Ser Cys Asp 115 120 125

Cys Leu Gln Gly Phe Gln Leu Thr His Ser Leu Gly Gly Gly Thr Gly 130 135 140

Ser Gly Met Gly Thr Leu Leu Ile Ser Lys Ile Arg Glu Glu Tyr Pro 145 150 155 160 Asp Arg Ile Met Asn Thr Phe Ser Val Val Pro Ser Pro Lys Val Ser 170 Asp Thr Val Val Glu Pro Tyr Asn Ala Thr Leu Ser Val His Gln Leu 185 Val Glu Asn Thr Asp Glu Thr Tyr Cys Ile Asp Asn Glu Ala Leu Tyr Asp Ile Cys Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp 215 Leu Asn His Leu Val Ser Ala Thr Met Ser Gly Val Thr Thr Cys Leu 230 Arg Phe Pro Gly Gln Leu Asn Ala Asp Leu Arg Lys Leu Ala Val Asn 245 Met Val Pro Phe Pro Arg Leu His Phe Phe Met Pro Gly Phe Ala Pro 265 260 Leu Thr Ser Arg Gly Ser Gln Gln Tyr Arg Ala Leu Thr Val Pro Glu 280 Leu Thr Gln Gln Met Phe Asp Ala Lys Asn Met Met Ala Ala Cys Asp 295 Pro Arg His Gly Arg Tyr Leu Thr Val Ala Ala Val Phe Arg Gly Arg 315 310 Met Ser Met Lys Glu Val Asp Glu Gln Met Leu Asn Val Gln Asn Lys 330 Asn Ser Ser Tyr Phe Val Glu Trp Ile Pro Asn Asn Val Lys Thr Ala 345 Val Cys Asp Ile Pro Pro Arg Gly Leu Lys Met Ser Ala Thr Phe Ile 360 Gly Asn Ser Thr Ala Ile Gln Glu Leu Phe Lys Arg Ile Ser Glu Gln 375 Phe Thr Ala Met Phe Arg Arg Lys Ala Phe Leu His Trp Tyr Thr Gly 395 390 Glu Gly Met Asp Glu Met Glu Phe Thr Glu Ala Glu Ser Asn Met Asn 410 Asp Leu Val Ser Glu Tyr Gln Gln Tyr Gln Asp Ala Thr Ala Glu Glu 420

Glu Gly Glu Phe Glu Glu Glu Ala Glu Glu Glu Val Ala

440

<210> 25

<211> 447

<212> PRT

<213> Neurospora crassa

<400> 25

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Gly Ala Ala Phe Trp Gln Thr Ile Ser Gly Glu His Gly Leu Asp Ala 20 25 30

Ser Gly Val Tyr Asn Gly Thr Ser Glu Leu Gln Leu Glu Arg Met Asn 35 40 45

Val Tyr Phe Asn Glu Ala Ser Gly Asn Lys Tyr Val Pro Arg Ala Val 50 55 60

Leu Val Asp Leu Glu Pro Gly Thr Met Asp Ala Val Arg Ala Gly Pro 65 70 75 80

Phe Gly Gln Leu Phe Arg Pro Asp Asn Phe Val Phe Gly Gln Ser Gly 85 90 95

Ala Gly Asn Asn Trp Ala Lys Gly His Tyr Thr Glu Gly Ala Glu Leu 100 105 110

Val Asp Gln Val Leu Asp Val Val Arg Arg Glu Ala Glu Gly Cys Asp 115 120 125

Cys Leu Gln Gly Phe Gln Ile Thr His Ser Leu Gly Gly Gly Thr Gly 130 135 140

Ala Gly Met Gly Thr Leu Leu Ile Ser Lys Ile Arg Glu Glu Phe Pro 145 150 155 160

Asp Arg Met Met Ala Thr Phe Ser Val Val Pro Ser Pro Lys Val Ser 165 170 175

Asp Thr Val Val Glu Pro Tyr Asn Ala Thr Leu Ser Val His Gln Leu 180 185 190

Val Glu Asn Ser Asp Glu Thr Phe Cys Ile Asp Asn Glu Ala Leu Tyr 195 . 200 205

Asp Ile Cys Met Arg Thr Leu Lys Leu Ser Asn Pro Ser Tyr Gly Asp 210 215 220

Leu Asn His Leu Val Ser Ala Val Met Ser Gly Val Thr Val Ser Leu 225 230 240

Arg Phe Pro Gly Gln Leu Asn Ser Asp Leu Arg Lys Leu Ala Val Asn 245 250 255

Met Val Pro Phe Pro Arg Leu His Phe Phe Met Val Gly Phe Ala Pro 260 265 270

Leu Thr Ser Arg Gly Ala His His Phe Arg Ala Val Ser Val Pro Glu Leu Thr Gln Gln Met Phe Asp Pro Lys Asn Met Met Ala Ala Ser Asp 295 Phe Arg Asn Gly Arg Tyr Leu Thr Cys Ser Ala Ile Phe Arg Gly Lys 310 Val Ser Met Lys Glu Val Glu Asp Gln Met Arg Asn Val Gln Asn Lys 330 Asn Ser Ser Tyr Phe Val Glu Trp Ile Pro Asn Asn Val Gln Thr Ala 345 Leu Cys Ser Ile Pro Pro Arg Gly Leu Lys Met Ser Ser Thr Phe Val Gly Asn Ser Thr Ala Ile Gln Glu Leu Phe Lys Arg Ile Gly Glu Gln Phe Thr Ala Met Phe Arg Arg Lys Ala Phe Leu His Trp Tyr Thr Gly 395 Glu Gly Met Asp Glu Met Glu Phe Thr Glu Ala Glu Ser Asn Met Asn 410 405 Asp Leu Val Ser Glu Tyr Gln Gln Tyr Gln Asp Ala Gly Val Asp Glu 420

<210> 26 <211> 447 <212> PRT <213> Aspergillus nidulans

435

Met Arg Glu lle Val His Leu Gln Thr Gly Gln Cys Gly Asn Gln lle Gly Ala Ala Phe 20 Thr Gly Asn Gly Asn Gly 30 Thr Gly Ala Tyr Asn Gly Ala Ser Gly Asn Gly Asn Gly Asn Gly Asn Tyr Asn Gly Ala Ser Gly Asn Leu Gln Leu Gly Arg Ala Val Leu Val Asp Leu Glu Pro Gly Thr Met Asp Cys Val Arg Ala Gly Pro

Glu Glu Glu Glu Tyr Glu Glu Glu Ala Pro Leu Glu Gly Glu Glu

Phe Gly Glu Leu Phe Arg Pro Asp Asn Phe Val Phe Gly Gln Ser Gly 85 90 95

Ala	Gly	Asn	Asn 100	Trp	Ala	Lys	Gly	His 105	Tyr	Thr	Glu	GLY	A1a 110	Glu	Leu
Val	Asp	Asn 115	Val	Val	Asp	Val	Val 120	Arg	Arg	Glu	Ala	Glu 125	Gly	Cys	Asp
Cys	Leu 130	Gln	Gly	Phe	Gln	Ile 135	Thr	His	Ser	Leu	Gly 140	Gly	Gly	Thr	Gly
Ala 145	Gly	Met	Gly	Thr	Leu 150	Leu	Ile	Ser	Lys	Ile 155	Arg	Glu	Glu	Phe	Pro 160
Asp	Arg	Met	Met	Ala 165	Thr	Phe	Ser	Val	Val 170	Pro	Ser	Pro	Lys	Val 175	Ser
Asp	Thr	Val	Val 180	Glu	Pro	Tyr	Asn	Ala 185	Thr	Leu	Ser	Val	His 190	Gln	Leu
Val	Glu	His 195	Ser	Asp	Glu	Thr	Phe 200	Cys	Ile	Asp	Asn	Glu 205	Ala	Leu	Туг
Asp	Ile 210	Cys	Met	Arg	Thr	Leu 215	Lys	Leu	Ser	Asn	Pro 220	Ser	Tyr	Gly	Asp
Leu 225	Asn	His	Leu	Val	Ser 230	Ala	Val	Met	Ser	Gly 235	Val	Thr	Thr	Суѕ	Leu 240
Arg	Phe	Pro	Gly	Gln 245	Leu	Asn	Ser	Asp	Leu 250	Arg	Lys	Trp	Ala	Val 255	Asn
Met	Val	Pro	Phe 260	Pro	Arg	Leu	His	Phe 265	Phe	Met	Val	Gly	Phe 270	Ala	Pro
Leu	Thr	Ser 275	Arg	Gly	Ala	Tyr	Ser 280	Phe	Arg	Ala	Val	Ser 285	Val	Pro	Glu
Leu	Thr 290	Gln	Gln	Met	Phe	Asp 295	Pro	Lys	Asn	Met	Met 300	Ala	Ala	Ser	Asp
Phe 305	Arg	Asn	Gly	Arg	Tyr 310	Leu	Thr	Cys	Ser	Ala 315	Ile	Phe	Arg	Gly	Lys 320
Val	Ser	Met	Lys	Glu 325	Val	Glu	Asp	Gln	Met 330	Arg	Asn	Ile	Gln	Ser 335	Lys
Asn	Gln	Ser	Tyr 340	Phe	Val	Glu	Trp	Ile 345	Pro	Asn	Asn	Ile	Gln 350	Ser	Ala
Leu	Cys	Ser 355	Ile	Pro	Pro	Arg	Gly 360	Leu	Lys	Met	Ser	Ser 365	Thr	Phe	Ile
Gly	Asn 370	Ser	Thr	Ser	Ile	Gln 375	Glu	Leu	Phe	Lys	Arg 380	Val	Gly	Asp	Glr
Phe 385	Thr	Ala	Met	Phe	Arg	Arg	Lys	Ala	Phe	Leu 395	His	Trp	Tyr	Thr	Gly 400

Glu Gly Met Asp Glu Met Glu Phe Thr Glu Ala Glu Ser Asn Met Asn

Asp Leu Val Ser Glu Tyr Gln Gln Tyr Gln Asp Ala Ser Ile Ser Glu

Gly Glu Glu Glu Tyr Ala Glu Glu Glu Ile Met Glu Gly Glu Glu 440

<210> 27

<211> 444

<212> PRT

<213> Achlya klebsiana

<400> 27

Met Arg Glu Leu Val His Ile Gln Gly Gln Cys Gly Asn Gln Ile

Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Val Asp Pro

Thr Gly Ser Tyr His Gly Asp Ser Asp Leu Gln Leu Glu Arg Ile Asn

Val Tyr Tyr Asn Glu Ala Thr Gly Thr Tyr Val Pro Arg Ala Ile Leu

Met Asp Leu Glu Pro Gly Thr Met Asp Ser Val Arg Ala Gly Pro Tyr

Gly Gln Leu Phe Arg Pro Asp Asn Phe Val Phe Gly Gln Thr Gly Ala

Gly Asn Asn Trp Ala Lys Gly His Tyr Thr Glu Gly Ala Glu Leu Ile

Asp Ser Val Leu Asp Val Val Arg Lys Glu Ala Glu Ser Cys Asp Cys

Leu Gln Gly Phe Gln Ile Thr His Ser Leu Gly Gly Gly Thr Gly Ser

Gly Met Gly Thr Leu Leu Ile Ser Lys Ile Arg Glu Glu Tyr Pro Asp

Arg Ile Met Cys Thr Tyr Ser Val Cys Pro Ser Pro Lys Val Ser Asp 170

Thr Val Val Glu Pro Tyr Asn Ala Thr Leu Ser Val His Gln Leu Val 185

Glu Asn Ala Asp Glu Val Met Cys Leu Asp Asn Glu Ala Leu Tyr Asp 200

Ile Cys Phe Arg Thr Leu Lys Leu Thr Asn Pro Thr Tyr Gly Asp Leu 210 215 220

Asn 225	His	Leu	Val	Cys	Ala 230	Ala	Met	Ser	Gly	Ile 235	Thr	Thr	Leu	Leu	Arg 240
Phe	Pro	Gly	Gln	Leu 245	Asn	Ser	Val	Leu	Lys 250	Leu	Ala	Val	Asn	Leu 255	Ile
Pro	Phe	Pro	Arg 260	Leu	His	Phe	Phe	Met 265	Ile	Gly	Phe	Ala	Pro 270	Leu	Thr
Ser	Arg	Gly 275	Ser	Gln	Gln	Tyr	Arg 280	Ala	Leu	Thr	Val	Pro 285	Glu	Leu	Thr
Gln	Gln 290	Gln	Phe	Asp	Ala	Lys 295	Asn	Met	Met	Cys	Ala 300	Ala	Asp	Pro	Arg
His 305	Gly	Arg	Tyr	Leu	Thr 310	Ala	Ala	Cys	Met	Phe 315	Arg	Gly	Arg	Met	Ser 320
Thr	Lys	Glu	Val	Asp 325	Glu	Gln	Met	Leu	Asn 330	Val	Gln	Asn	Lys	Asn 335	Ser
Ser	Tyr	Phe	Val 340	Glu	Trp	Ile	Pro	Asn 345	Asn	Ile	Lys	Ala	Ser 350	Val	Cys
Asp	Ile	Pro 355	Pro	Lys	Gly	Leu	Lys 360	Met	Ser	Thr	Thr	Phe 365	Ile	Gly	Asn
Ser	Thr 370	Ala	Ile	Gln	Glu	Met 375	Phe	Lys	Arg	Val	Ser 380	Glu	Gln	Phe	Thr
Ala 385	Met	Phe	Arg	Arg	Lys 390	Ala	Phe	Leu	His	Trp 395	Tyr	Thr	Gly	Glu	Gly 400
Met	Asp	Glu	Met	Glu 405	Phe	Thr	Glu	Ala	Glu 410	Ser	Asn	Met	Asn	Asp 415	Leu
Val	Ser	Glu	Tyr 420	Gln	Gln	Tyr	Gln	Asp 425	Ala	Thr	Ala	Glu	Glu 430	Glu	Gly
Glu	Phe	Asp 435	Glu	Asp	Glu	Glu	Met 440	Asp	Glu	Met	Met				

<210> 28

<211> 15

<212> PRT

<213> Pestalotiopsis microspora

<220>

<223> motif conserved in P. ultimum, H. sapiens, N. crassa, A. nidulans, and A. klebsiana

<400> 28

Ala Ile Leu Val Asp Leu Glu Pro Gly Thr Met Asp Ser Val Arg
1 5 10 15

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<210> 29
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<213> Pestalotiopsis microspora
<220>
<223> motif conserved in P. ultimum, H. sapiens, N.
      crassa, A. nidulans, and A. klebsiana
<400> 29
Ala Val Leu Val Asp Leu Glu Pro Gly Thr Met Asp Ser Val Arg
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<220>
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      crassa, A. nidulans, and A. klebsiana
<400> 30
Gly Gly Gly Thr Gly Ser Gly
<210> 31
<211> 4
<212> PRT
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<220>
<223> motif conserved in P. ultimum, H. sapiens, N.
      crassa, A. nidulans, and A. klebsiana
<400> 31
Asp Asn Glu Ala
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<210> 32
<211> 4
<212> PRT
<213> Pestalotiopsis microspora
<220>
<223> motif conserved in P. ultimum, H. sapiens, N.
      crassa, A. nidulans and A. klebsiana
<400> 32
Met Arg Glu Ile
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<210> 33
<211> 4
<212> PRT
<213> Pestalotiopsis microspora
<220>
<223> motif conserved in P. ultimum, H. sapiens, N.
     crassa, A. nidulans, and A. klebsiana
<400> 33
Met Arg Glu Leu
<210> 34
<211> 31
<212> PRT
<213> Sus scrofa
<400> 34
Met Arg Glu Ile Val His Ile Gln Ala Gly Gln Cys Gly Asn Gln Ile
Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Ile Asp
             20
<210> 35
<211> 20
<212> PRT
<213> Sus scrofa
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
Leu Val Ser Ala
<210> 36
<211> 31
<212> PRT
<213> Homo sapiens
<400> 36
Met Arg Glu Ile Val His Leu Gln Ala Gly Gln Cys Gly Asn Gln Ile
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Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Ile Asp
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<210> 37
<211> 20
<212> PRT
<213> Homo sapiens
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<400> 37
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
Leu Val Ser Ala
<210> 38
<211> 31
<212> PRT
<213> Drosophila melanogaster
<400> 38
Met Arg Glu Ile Val His Ile Gln Ala Gly Gln Cys Gly Asn Gln Ile
Gly Ala Lys Phe Trp Glu Ile Ile Ser Asp Glu His Gly Ile Asp
<210> 39
<211> 20
<212> PRT
<213> Drosophila melanogaster
<400> 39
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
                                     10
Leu Val Ser Leu
<210> 40
<211> 31
<212> PRT
<213> Xenopus laevis
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Met Arg Glu Ile Val His Leu Gln Ala Gly Gln Cys Gly Asn Gln Ile
Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Ile Asp
<210> 41
<211> 20
<212> PRT
<213> Xenopus laevis
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
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Leu Val Ser Ala

Leu Val Ser Ala

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<210> 42
<211> 31
<212> PRT
<213> Tetrahymena thermophila
<400> 42
Met Arg Glu Ile Val His Ile Gln Gly Gly Gln Cys Gly Asn Gln Ile
Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Ile Asp
<210> 43
<211> 20
<212> PRT
<213> Tetrahymena thermophila
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
Leu Val Ser Ala
<210> 44
<211> 31
<212> PRT
<213> Physarum polycephalum
<400> 44
Met Arg Glu Ile Val His Ile Gln Ala Gly Gln Cys Gly Asn Gln Ile
Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Ile Asp
             20
<210> 45
<211> 20
<212> PRT
<213> Physarum polycephalum
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
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<210> 46
<211> 31
<212> PRT
<213> Pythium ultimum
<400> 46
Met Arg Glu Leu Val His Ile Gln Gly Gly Gln Cys Gly Asn Gln Ile
Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Val Asp
<210> 47
<211> 20
<212> PRT
<213> Pythium ultimum
<400> 47
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
                  5
Leu Val Cys Ala
<210> 48
<211> 31
<212> PRT
<213> Phytophthora cinnamomi
<400> 48
Met Arg Glu Leu Val His Ile Gln Gly Gly Gln Cys Gly Asn Gln Ile
Gly Ala Lys Phe Trp Glu Val Val Ser Asp Glu His Gly Val Asp
<210> 49
<211> 20
<212> PRT
<213> Phytophthora cinnamomi
<400> 49
Phe Arg Thr Leu Lys Leu Thr Thr Pro Thr Tyr Gly Asp Leu Asn His
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Leu Val Cys Ala
<210> 50
<211> 31
<212> PRT
<213> Achlya klebsiana
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<400> 50

30

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Gly Ala Lys Phe Trp Glu Val Ile Ser Asp Glu His Gly Val Asp
<210> 51
<211> 20
<212> PRT
<213> Achlya klebsiana
<400> 51
Phe Arg Thr Leu Lys Leu Thr Asn Pro Thr Tyr Gly Asp Leu Asn His
Leu Val Cys Ala
<210> 52
<211> 31
<212> PRT
<213> Pestalotiopsis microspora
Met Arg Glu Ile Val His Leu Gln Thr Gly Gln Cys Gly Asn Gln Ile
Gly Ala Ala Phe Trp Gln Thr Ile Ser Gly Glu His Gly Leu Asp
<210> 53
<211> 20
<212> PRT
<213> Pestalotiopsis microspora
<400> 53
Met Arg Thr Leu Lys Leu Ser Asn Pro Ser Tyr Gly Asp Leu Asn His
Leu Val Ser Ala
<210> 54
<211> 31
<212> PRT
<213> Aspergillus nidulans
Met Arg Glu Ile Val His Leu Gln Thr Gly Gln Cys Gly Asn Gln Ile
                                     10
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